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(57) Abstract

Compounds containing two aromatic systems covalently linked through a linker containing one or more atoms, or "linker" defined as including a covalent bond per are so as to space the aromatic systems as a distance 15-15Å, are effective in treating contains associated with bone deficits. The compounds can be administered to vertebrate subjects alone or in combination with additional agents that promote bone growth or that inhibit bone recoprofton. They can be screened for activity prior to administration by assessing their like the transcription of a reporter gene coupled to a promoter associated with a bone morphogenetic protein and/or their ability to effect calvarial growth in model animal systems.

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COMPOSITIONS AND METHODS FOR TREATING BONE DEFICIT CONDITIONS

Technical Field

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The invention relates to compositions and methods for use in limiting undesired bone loss in a vertebrate at risk of such bone loss, in treating conditions that are characterized by undesired bone loss or by the need for bone growth, in treating fractures, and in treating cartilage disorders. More specifically, the invention concerns the use of specific classes of compounds identified or characterized by a high throughput screening assay.

Background Art

Bone is not a static tissue. It is subject to constant breakdown and resynthesis in a complex process mediated by osteoblasts, which produce new bone, and osteoclasts, which destroy bone. The activities of these cells are regulated by a large number of cytokines and growth factors, many of which have now been identified and cloned. Mundy has described the current knowledge related to these factors (Mundy, G.R. Clin Orthop 324:24-28, 1996; Mundy, G.R. J Bone Miner Res 8:S505-10, 1993)

Although there is a great deal of information available on the factors which influence the breakdown and resorption of bone, information on growth factors which stimulate the formation of new bone is more limited. Investigators have searched for sources of such activities, and have found that bone tissue itself is a storehouse for factors which have the capacity for stimulating bone cells. Thus, extracts of bovine bone tissue obtained from slaughterhouses contain not only structural proteins which are responsible for maintaining the structural integrity of bone, but also biologically active bone growth factors which can stimulate bone cells to proliferate. Among these latter factors are transforming growth factor β , the heparin-binding growth factors (acidic and basic fibroblast growth factor), the insulin-like growth factor I and insulin-like growth factor II), and a recently described family of

proteins called bone morphogenetic proteins (BMPs). All of these growth factors have effects on other types of cells, as well as on bone cells.

The BMPs are novel factors in the extended transforming growth factor B superfamily. They were first identified by Wozney J. et al. Science (1988) 242:1528-34, using gene cloning techniques, following earlier descriptions characterizing the biological activity in extracts of demineralized bone (Urist M. Science (1965) 150:893-99). Recombinant BMP2 and BMP4 can induce new bone formation when they are injected locally into the subcutaneous tissues of rats (Wozney J. Molec Reprod Dev (1992) 32:160-67). These factors are expressed by normal osteoblasts as they differentiate, and have been shown to stimulate osteoblast differentiation and bone nodule formation in vitro as well as bone formation in vivo (Harris S. et al. J. Bone Miner Res (1994) 9:855-63). This latter property suggests potential usefulness as therapeutic agents in diseases which result in bone loss.

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The cells which are responsible for forming bone are osteoblasts. As osteoblasts differentiate from precursors to mature bone-forming cells, they express and secrete a number of enzymes and structural proteins of the bone matrix, including Type-1 collagen, osteocalcin, osteopontin and alkaline phosphatase (Stein G. et al. Curr Opin Cell Biol (1990) 2:1018-27; Harris S. et al. (1994), supra). They also synthesize a number of growth regulatory peptides which are stored in the bone matrix, and are presumably responsible for normal bone formation. These growth regulatory peptides include the BMPs (Harris S. et al. (1994), supra). In studies of primary cultures of fetal rat calvarial osteoblasts, BMPs 1, 2, 3, 4, and 6 are expressed by cultured cells prior to the formation of mineralized bone nodules (Harris S. et al. (1994), supra). Like alkaline phosphatase, osteocalcin and osteopontin, the BMPs are expressed by cultured osteoblasts as they proliferate and differentiate.

Although the BMPs are potent stimulators of bone formation in vitro and in vivo, there are disadvantages to their use as therapeutic agents to enhance bone healing. Receptors for the bone morphogenetic proteins have been identified in many tissues, and the BMPs themselves are expressed in a large variety of tissues in specific temporal and spatial patterns. This suggests that BMPs may have effects on many

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tissues other than bone, potentially limiting their usefulness as therapeutic agents when administered systemically. Moreover, since they are peptides, they would have to be administered by injection. These disadvantages impose severe limitations to the development of BMPs as therapeutic agents.

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There is a plethora of conditions which are characterized by the need to enhance bone formation. Perhaps the most obvious is the case of bone fractures, where it would be desirable to stimulate bone growth and to hasten and complete bone repair. Agents that enhance bone formation would also be useful in facial reconstruction procedures. Other bone deficit conditions include bone segmental defects, periodontal disease, metastatic bone disease, osteolytic bone disease and conditions where connective tissue repair would be beneficial, such as healing or regeneration of cartilage defects or injury. Also of great significance is the chronic condition of osteoporosis, including age-related osteoporosis and osteoporosis associated with postmenopausal hormone status. Other conditions characterized by the need for bone growth include primary and secondary hyperparathyroidism, disuse osteoporosis, diabetes-related osteoporosis, and glucocorticoid-related osteoporosis. In addition, or alternatively, the compounds of the present invention may modulate metabolism, proliferation and/or differentiation of normal or aberrant cells or tissues.

There are currently no satisfactory pharmaceutical approaches to managing any of these conditions. Bone fractures are still treated exclusively using casts, braces, anchoring devices and other strictly mechanical means. Further bone deterioration associated with postmenopausal osteoporosis has been decreased or prevented with estrogens or bisphosphonates.

US Patent 5, 280, 040 discloses a class of compounds which are 3, 4-diaryl chromans. These compounds can be considered derivatives of 2,3,4 triphenyl butanol, where the hydroxy at the 1-position forms an ether with the ortho position of the phenyl group substituted at the 4-position of the butanol. The parent 3,4-diaryl chromans do not contain nitrogen atoms in the aromatic moieties or their linkers. A preferred compound, centchroman, contains a nitrogen substituent only in one of the

substituents on a phenyl moiety. These compounds are disclosed in the '040 patent as useful in the treatment of osteoporosis.

In addition, the PCT application WO97/15308 published 1 May 1997 describes a number of classes of compounds that are active in the screening assay described below and are useful in treating bone disorders. These compounds, generically, are of the formulae

$$R_{m}^{a}$$
 Z
 Z
 $L-Ar^{2}$

wherein R* is a non-interfering substituent;

m is an integer of 0-4;

each dotted line represents an optional π -bond:

each Z is independently N, NR, O, S, CR or CR2, where each R is

X is O, S, SO or SO2;

L is a flexible linker: and

Ar² is a substituted or unsubstituted 6-membered aromatic ring; or:

wherein R2 is a non-interfering substituent;

n is an integer of 0 and 5;

L is a flexible linker which does not contain nitrogen or is a constrained linker;

and

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 Ar^2 is a substituted or unsubstituted phenyl or a substituted or unsubstituted naphthyl.

There remains a need for additional compositions which can ameliorate the effects of abnormalities in bone formation or resorption. The present invention

expands the repertoire of compounds useful for limiting or treating bone deficit conditions, and for other uses that should be apparent to those skilled in the art from the teachings herein.

5 Disclosure of the Invention

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The invention provides compounds that can be administered as ordinary pharmaceuticals and have the metabolic effect of enhancing bone growth or inhibiting resorption. The compounds of the invention can be identified using an assay for their ability to activate control elements associated with bone anabolic factors. Thus, the invention is directed to methods and compositions for treating bone disorders, which methods and compositions use, as active ingredients, compounds wherein two aromatic systems are coupled so as to be spaced apart from each other by about 1.5 to about 15 Angstroms. The thus-linked systems (including the linker coupling them) preferably include at least one nitrogen atom.

Therefore, the compounds useful in the invention can be described as having the formula Ar¹-linker-Ar², wherein each of Ar¹ and Ar² is independently an aromatic system and the linker portion of the formula spaces Ar¹ and Ar² apart by a distance of approximately 1.5-15 Angstroms. Ar¹, Ar² and the linker may optionally be substituted with non interfering substituents. In the useful compounds, there is preferably at least one nitrogen atom in either Ar¹, Ar² and/or the linker, independent of any substituents thereon. Preferably, the compounds of the invention contain at least one additional heteroatom selected from the group consisting of N, S and O, independent of any substituent.

Thus, in one aspect, the invention is directed to a method to treat a condition in a vertebrate animal characterized by a deficiency in, or need for, bone growth replacement and/or an undesirable level of bone resorption, which method comprises administering to a vertebrate subject in need of such treatment an effective amount of certain compounds of the formula:

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wherein each of Ar1 and Ar2 is independently substituted or unsubstituted phenyl, substituted or unsubstituted naphthyl, a substituted or unsubstituted aromatic system containing a 6-membered heterocycle, or a substituted or unsubstituted aromatic system containing a 5-membered heterocycle; and

L is a linker that provides spacing of 1.5-15Å.

In other aspects, the invention relates to pharmaceutical compositions for use in the method, and to the compounds for use in preparing a medicament for use in the method.

10 Brief Description of the Drawings

Figure 1 gives a schematic representation of the compounds used as active ingredients in the methods and compositions of the invention.

Figure 2 shows the dose response curve for a positive control compound. designated 59-0008.

Figures 3 and 4 show illustrative compounds of the invention and the results obtained with them in an in vitro test for stimulation of bone growth.

Figures 5A, 5B and 5C show structures and results of a screening assay for a group of compounds which varies the parameters of lead compound 59-0072.

Figures 6A, 6B and 6C show structures and results of a screening assay for a group of compounds which varies the parameters of lead compound 50-0197.

Figure 7 shows structures and results of a screening assay for a group of compounds which varies the parameters of lead compound 59-0145.

Figures 8A, 8B and 8C show structures and results of a screening assay for a group of compounds which varies the parameters of lead compound 59-0045.

25 Figure 9 shows the results in an ex vivo calvarial assay for various compunds of the invention

Figure 10 shows the increase in bone volume effected by subcutaneous administration of compound 59-0145 in the OVX in vivo assay.

Figure 11 is a graphical representation of percent increase in trabecular bone in ovariectomized rats treated with compound 59-0145

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Figure 12 presents graphs showing results of qCT and bone histomorphometri and serum osteocalcin levels in rats treated with compound 59-0145.

Figure 13 (41 pages) is a list of compounds used in screening for bone morphogenic activity according to the screening assay set forth herein.

Modes of Carrying Out the Invention

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A rapid throughput screening test for compounds capable of stimulating expression of a reporter gene linked to a BMP promoter (a surrogate for the production of bone morphogenetic factors that are endogenously produced) is described in WO96/38590 published 5 December 1996, the contents of which are incorporated herein by reference. This assay is also described as a portion of a study of immortalized murine osteoblasts (derived from a mouse expressing a transgene composed of a BMP2 promoter driving expression of T-antigen) in Ghosh-Choudhery, N. et al. Endocrinology (1996) 137:331-39. In this study, the immortalized cells were stably transfected with a plasmid containing a luciferase reporter gene driven by a mouse BMP2 promoter (-2736/114 bp), and responded in a dose-dependent manner to recombinant human BMP2.

Briefly, the assay utilizes cells transformed permanently or transiently with constructs in which the promoter of a bone morphogenetic protein, specifically BMP2 or BMP4, is coupled to a reporter gene, typically luciferase. These transformed cells are then evaluated for the production of the reporter gene product; compounds that activate the BMP promoter will drive production of the reporter protein, which can be readily assayed. Over 40,000 compounds have been subjected to this rapid screening technique, and only a very small percentage are able to elicit a level of production of luciferase 5-fold greater than that produced by vehicle. Compounds that activate the BMP promoter share certain structural characteristics not present in inactive compounds. The active compounds ("BMP promoter-active compounds") or "active compounds") are useful in promoting bone or cartilage growth, and thus in the treatment of vertebrates in need of bone or cartilage growth.

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BMP promoter-active compounds can be examined in a variety of other assays that test specificity and toxicity. For instance, nonBMP promoters or response elements can be linked to a reporter gene and inserted into an appropriate host cell. Cytotoxicity can be determined by visual or microscopic examination of BMP promoter- and/or nonBMP promoter-reporter gene-containing cells, for instance. Alternatively, nucleic acid and/or protein synthesis by the cells can be monitored. For in vivo assays, tissues may be removed and examined visually or microscopically, and optionally examined in conjunction with dyes or stains that facilitate histologic examination. In assessing in vivo assay results, it may also be useful to examine biodistribution of the test compound, using conventional medicinal chemistry/animal model techniques.

As used herein, "limit" or "limiting" and "treat" or "treatment" are interchangeable terms. The terms include a postponement of development of bone deficit symptoms and/or a reduction in the severity of such symptoms that will or are expected to develop. The terms further include ameliorating existing bone or cartilage deficit symptoms, preventing additional symptoms, ameliorating or preventing the underlying metabolic causes of symptoms, preventing or reversing bone resorption and/or encouraging bone growth. Thus, the terms denote that a beneficial result has been conferred on a vertebrate subject with a cartilage, bone or skeletal deficit, or with the potential to develop such deficit.

By "bone deficit" is meant an imbalance in the ratio of bone formation to bone resorption, such that, if unmodified, the subject will exhibit less bone than desirable, or the subject's bones will be less intact and coherent than desired. Bone deficit may also result from fracture, from surgical intervention or from dental or periodontal disease. By "cartilage defect" is meant damaged cartilage, less cartilage than desired, or cartilage that is less intact and coherent than desired.

Representative uses of the compounds of the present invention include: repair of bone defects and deficiencies, such as those occurring in closed, open and nonunion fractures, prophylactic use in closed and open fracture reduction; promotion of bone healing in plastic surgery, stimulation of bone ingrowth into noncemented prosthetic

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joints and dental implants; elevation of peak bone mass in premenopausal women; treatment of growth deficiencies; treatment of peridontal disease and defects, and other tooth repair processes; increase in bone formation during distraction osteogenesis; and treatment of other skeletal disorders, such as age-related osteoporosis, postmenopausal osteoporosis, glucocorticoid-induced osteoporosis or disuse osteoporosis and arthritis. The compounds of the present invention can also be useful in repair of congenital, trauma-induced or surgical resection of bone (for instance, for cancer treatment), and in cosmetic surgery. Further, the compounds of the present invention can be used for limiting or treating cartilage defects or disorders, and may be useful in wound healing or tissue repair.

Bone or cartilage deficit or defect can be treated in vertebrate subjects by administering compounds of the invention which have been identified through suitable screening assays and which exhibit certain structural characteristics. The compositions of the invention may be administered systemically or locally. For systemic use, the compounds herein are formulated for parenteral (e.g., intravenous, subcutaneous, intramuscular, intraperitoneal, intranasal or transdermal) or enteral (e.g., oral or rectal) delivery according to conventional methods. Intravenous administration will be by a series of injections or by continuous infusion over an extended period. Administration by injection or other routes of discretely spaced administration will generally be performed at intervals ranging from weekly to once to three times daily. Alternatively, the compounds disclosed herein may be administered in a cyclical manner (administration of disclosed compound; followed by no administration; followed by administration of disclosed compound, and the like). Treatment will continue until the desired outcome is achieved. In general, pharmaceutical formulations will include a compound of the present invention in combination with a pharmaceutically acceptable vehicle, such as saline, buffered saline, 5% dextrose in water, borate-buffered saline containing trace metals or the like. Formulations may further include one or more excipients, preservatives, solubilizers, buffering agents, albumin to prevent protein loss on vial surfaces, lubricants, fillers, stabilizers, etc. Methods of formulation are well known in the art and are disclosed, for example, in Remington's Pharmaceutical

Sciences, Gennaro, ed., Mack Publishing Co., Easton PA, 1990, which is incorporated herein by reference. Pharmaceutical compositions for use within the present invention can be in the form of sterile, nonpyrogenic liquid solutions or suspensions, coated capsules, suppositories, lyophilized powders, transdermal patches or other forms known in the art. Local administration may be by injection at the site of injury or defect, or by insertion or attachment of a solid carrier at the site, or by direct, topical application of a viscous liquid. For local administration, the delivery vehicle preferably provides a matrix for the growing bone or cartilage, and more preferably is a vehicle that can be absorbed by the subject without adverse effects.

Delivery of compounds herein to wound sites may be enhanced by the use of controlled-release compositions, such as those described in WIPO publication WO 93/20859, which is incorporated herein by reference in its entirety. Films of this type are particularly useful as coatings for prosthetic devices and surgical implants. The films may, for example, be wrapped around the outer surfaces of surgical screws, rods, pins, plates and the like. Implantable devices of this type are routinely used in orthopedic surgery. The films can also be used to coat bone filling materials, such as hydroxyapatite blocks, demineralized bone matrix plugs, collagen matrices and the like. In general, a film or device as described herein is applied to the bone at the fracture site. Application is generally by implantation into the bone or attachment to the surface using standard surgical procedures.

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In addition to the copolymers and carriers noted above, the biodegradable films and matrices may include other active or inert components. Of particular interest are those agents that promote tissue growth or infiltration, such as growth factors.

Exemplary growth factors for this purpose include epidermal growth factor (EGF), fibroblast growth factor (FGF), platelet-derived growth factor (PDGF), transforming growth factors (TGFs), parathyroid hormone (PTH), leukemia inhibitory factor (LIF), and insulin-like growth factors (IGFs). Agents that promote bone growth, such as bone morphogenetic proteins (U.S. Patent No. 4,761,471; PCT Publication WO 90/11366), osteogenin (Sampath et al. Proc. Natl. Acad. Sci. USA (1987) 84:7109-13) and NaF (Tencer et al. J. Biomed. Mat. Res. (1989) 23: 571-89) are also preferred.

Biodegradable films or matrices include calcium sulfate, tricalcium phosphate, hydroxyapatite, polylactic acid, polyanhydrides, bone or dermal collagen, pure proteins, extracellular matrix components and combinations thereof. Such biodegradable materials may be used in combination with nonbiodegradable materials, to provide desired mechanical, cosmetic or tissue or matrix interface properties.

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Alternative methods for delivery of compounds of the present invention include use of ALZET osmotic minipumps (Alza Corp., Palo Alto, CA); sustained release matrix materials such as those disclosed in Wang et al. (PCT Publication WO 90/11366), electrically charged dextran beads, as disclosed in Bao et al. (PCT Publication WO 92/03125); collagen-based delivery systems, for example, as disclosed in Ksander et al. Arm. Surg. (1990) 211(3):288-94; methylcellulose gel systems, as disclosed in Beck et al. J. Bone Min. Res. (1991) 6(11):1257-65; and alginate-based systems, as disclosed in Edelman et al. Biomaterials (1991) 12:619-26. Other methods well known in the art for sustained local delivery in bone include porous coated metal protheses that can be impregnated and solid plastic rods with therapeutic compositions incorporated within them

The compounds of the present invention may also be used in conjunction with agents that inhibit bone resorption. Antiresorptive agents, such as estrogen, bisphosphonates and calcitonin, are preferred for this purpose. More specifically, the compounds disclosed herein may be administered for a period of time (for instance, months to years) sufficient to obtain correction of a bone deficit condition. Once the bone deficit condition has been corrected, the vertebrate can be administered an anti-resorptive compound to maintain the corrected bone condition. Alternatively, the compounds disclosed herein may be administered with an anti-resorptive compound in a cyclical manner (administration of disclosed compound, followed by anti-resorptive, followed by disclosed compound, and the like).

In additional formulations, conventional preparations such as those described below may be used.

Aqueous suspensions may contain the active ingredient in admixture with pharmacologically acceptable excipients, comprising suspending agents, such as methyl cellulose; and wetting agents, such as lecithin, lysolethicin or long-chain fatty alcohols.

The said aqueous suspensions may also contain preservatives, coloring agents,
flavoring agents and sweetening agents in accordance with industry standards.

Preparations for topical and local application comprise aerosol sprays, lotions, gels and ointments in pharmaceutically appropriate vehicles which may comprise lower aliphatic alcohols, polyglycols such as glycerol, polyethylene glycol, esters of fatty acids, oils and fats, and silicones. The preparations may further comprise antioxidants, such as ascorbic acid or tocopherol, and preservatives, such as p-hydroxybenzoic acid esters.

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Parenteral preparations comprise particularly sterile or sterilized products.

Injectable compositions may be provided containing the active compound and any of the well known injectable carriers. These may contain salts for regulating the osmotic pressure.

If desired, the osteogenic agents can be incorporated into liposomes by any of the reported methods of preparing liposomes for use in treating various pathogenic conditions. The present compositions may utilize the compounds noted above incorporated in liposomes in order to direct these compounds to macrophages, monocytes, other cells and tissues and organs which take up the liposomal composition. The liposome-incorporated compounds of the invention can be utilized by parenteral administration, to allow for the efficacious use of lower doses of the compounds. Ligands may also be incorporated to further focus the specificity of the liposomes.

Suitable conventional methods of liposome preparation include, but are not limited to, those disclosed by Bangham, A.D. et al. J Mol Biol (1965) 23:238-252, Olson, F. et al. Biochim Biophys Acta (1979) 557:9-23, Szoka, F. et al. Proc Natl Acad Sci USA (1978) 75:4194-4198, Mayhew, E. et al. (1984) 775:169-175, Kim, S. et al. Biochim Biophys Acta (1983) 728:339:348, and Mayer, et al. Biochim Biophys Acta (1986) 858:161-168.

The liposomes may be made from the present compounds in combination with

30 any of the conventional synthetic or natural phospholipid liposome materials including

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phospholipids from natural sources such as egg, plant or animal sources such as phosphatidylcholine, phosphatidylethanolamine, phosphatidylglycerol, sphingomyelin, phosphatidylserine, or phosphatidylinositol. Synthetic phospholipids that may also be used, include, but are not limited to: dimyristoylphosphatidylcholine,

5 dioleoylphosphatidylcholine, dipalmitoylphosphatidylcholine and distearoylphosphatidylcholine, and the corresponding synthetic phosphatidylethanolamines and phosphatidylglycerols. Cholesterol or other sterols, cholesterol hemisuccinate, glycolipids, cerebrosides, fatty acids, gangliosides, sphingolipids, 1,2-bis(oleoyloxy)-3-(trimethyl ammonio) propane (DOTAP), N-[1-10 (2 3-dioleoyl) provyl-N N-trimethylammonium chloride (DOTAM), and other.

(2,3-dioleoyl) propyl-N,N,N-trimethylammonium chloride (DOTMA), and other cationic lipids may be incorporated into the liposomes, as is known to those skilled in the art. The relative amounts of phospholipid and additives used in the liposomes may be varied if desired. The preferred ranges are from about 60 to 90 mole percent of the phospholipid, cholesterol, cholesterol hemisuccinate, fatty acids or cationic lipids may be used in amounts ranging from 0 to 50 mole percent. The amounts of the present compounds incorporated into the lipid layer of liposomes can be varied with the concentration of their lipids ranging from about 0.01 to about 50 mole percent.

Using conventional methods, approximately 20 to 30% of the compound present in solution can be entrapped in liposomes; thus, approximately 70 to 80% of the active compound is wasted. In contrast, where the compound is incorporated into liposomes, virtually all of the compound is incorporated into the liposome, and essentially none of the active compound is wasted.

The liposomes with the above formulations may be made still more specific for their intended targets with the incorporation of monoclonal antibodies or other ligands specific for a target. For example, monoclonal antibodies to the BMP receptor may be incorporated into the liposome by linkage to phosphatidylethanolamine (PE) incorporated into the liposome by the method of Leserman, L. et al. Nature (1980) 288:602-604.

Veterinary uses of the disclosed compounds are also contemplated. Such uses would include limitation or treatment of bone or cartilage deficits or defects in

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domestic animals, livestock and thoroughbred horses. The compounds described herein can also modify a target tissue or organ environment, so as to attract boneforming cells to an environment in need of such cells.

The compounds of the present invention may also be used to stimulate growth of bone-forming cells or their precursors, or to induce differentiation of bone-forming cell precursors, either in vitro or ex vivo. As used herein, the term "precursor cell" refers to a cell that is committed to a differentiation pathway, but that generally does not express markers or function as a mature, fully differentiated cell. As used herein. the term "mesenchymal cells" or "mesenchymal stem cells" refers to pluripotent progenitor cells that are capable of dividing many times, and whose progeny will give rise to skeletal tissues, including cartilage, bone, tendon, ligament, marrow stroma and connective tissue (see A. Caplan J. Orthop. Res. (1991) 9:641-50). As used herein, the term "osteogenic cells" includes osteoblasts and osteoblast precursor cells. More particularly, the disclosed compounds are useful for stimulating a cell population containing marrow mesenchymal cells, thereby increasing the number of osteogenic cells in that cell population. In a preferred method, hematopoietic cells are removed from the cell population, either before or after stimulation with the disclosed compounds. Through practice of such methods, osteogenic cells may be expanded. The expanded osteogenic cells can be infused (or reinfused) into a vertebrate subject in need thereof. For instance, a subject's own mesenchymal stem cells can be exposed to compounds of the present invention ex vivo, and the resultant osteogenic cells could be infused or directed to a desired site within the subject, where further proliferation and/or differentiation of the osteogenic cells can occur without immunorejection. Alternatively, the cell population exposed to the disclosed compounds may be immortalized human fetal osteoblastic or osteogenic cells. If such cells are infused or implanted in a vertebrate subject, it may be advantageous to "immunoprotect" these nonself cells, or to immunosuppress (preferably locally) the recipient to enhance transplantation and bone or cartilage repair.

Within the present invention, an "effective amount" of a composition is that amount which produces a statistically significant effect. For example, an "effective

amount" for therapeutic uses is the amount of the composition comprising an active compound herein required to provide a clinically significant increase in healing rates in fracture repair; reversal of bone loss in osteoporosis; reversal of cartilage defects or disorders; prevention or delay of onset of osteoporosis; stimulation and/or augmentation of bone formation in fracture nonunions and distraction osteogenesis: increase and/or acceleration of bone growth into prosthetic devices; and repair of dental defects. Such effective amounts will be determined using routine optimization techniques and are dependent on the particular condition to be treated, the condition of the patient, the route of administration, the formulation, and the judgment of the practitioner and other factors evident to those skilled in the art. The dosage required for the compounds of the invention (for example, in osteoporosis where an increase in bone formation is desired) is manifested as a statistically significant difference in bone mass between treatment and control groups. This difference in bone mass may be seen, for example, as a 5-20% or more increase in bone mass in the treatment group. Other measurements of clinically significant increases in healing may include, for example, tests for breaking strength and tension, breaking strength and torsion, 4-point bending, increased connectivity in bone biopsies and other biomechanical tests well known to those skilled in the art. General guidance for treatment regimens is obtained from experiments carried out in animal models of the disease of interest

The dosage of the compounds of the invention will vary according to the extent and severity of the need for treatment, the activity of the administered compound, the general health of the subject, and other considerations well known to the skilled artisan. Generally, they can be administered to a typical human on a daily basis on an oral dose of about 0.1 mg/kg-1000 mg/kg, and more preferably from about 1 mg/kg to about 200 mg/kg. The parenteral dose will appropriately be 20-100% of the oral dose.

Screening Assays

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The osteogenic activity of the compounds used in the methods of the invention can be verified using *in vitro* screening techniques, such as the assessment of

transcription of a reporter gene coupled to a bone morphogenetic protein-associated

Technique for Neonatal Mouse Calvarial Assay (In vitro)

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promoter, as described above, or in alternative assays such as the following:

This assay is similar to that described by Gowen M. & Mundy G. *J Immunol* (1986) 136:2478-82. Briefly, four days after birth, the front and parietal bones of ICR Swiss white mouse pups are removed by microdissection and split along the sagittal suture. The bones are incubated in BGJb medium (Irvine Scientific, Santa Ana, CA) plus 0.02% (or lower concentration) β -methylcyclodextrin, wherein the medium also contains test or control substances, at 37°C in a humidified atmosphere of 5% CO₂ and 95% air for 96 hours.

Following this, the bones are removed from the incubation media and fixed in 10% buffered formalin for 24-48 hours, decalcified in 14% EDTA for 1 week, processed through graded alcohols; and embedded in paraffin wax. Three µm sections of the calvaria are prepared. Representative sections are selected for histomorphometric assessment of bone formation and bone resorption. Bone changes are measured on sections cut 200 µm apart. Osteoblasts and osteoclasts are identified by their distinctive morphology.

Other auxillary assays can be used as controls to determine nonBMP promotermediated effects of test compounds. For example, mitogenic activity can be measured
using screening assays featuring a serum-response element (SRE) as a promoter and a
luciferase reporter gene. More specifically, these screening assays can detect signalling
through SRE-mediated pathways, such as the protein kinase C pathway. For instance,
an osteoblast activator SRE-luciferase screen and an insulin mimetic SRE-luciferase
screen are useful for this purpose. Similarly, test compound stimulation of cAMP
response element (CRE)-mediated pathways can also be assayed. For instance, cells
transfected with receptors for PTH and calcitonin (two bone-active agents) can be
used in CRE-luciferase screens to detect elevated cAMP levels. Thus, the BMP
promoter specificity of a test compound can be examined through use of these types of
auxillary assays.

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In vivo Assay of Effects of Compounds on Murine Calvarial Bone Growth Male ICR Swiss white mice, aged 4-6 weeks and weighing 13-26 gm, are employed, using 4-5 mice per group. The calvarial bone growth assay is performed as described in PCT application WO 95/24211. Briefly, the test compound or appropriate control vehicle is injected into the subcutaneous tissue over the right calvaria of normal mice. Typically, the control vehicle is the vehicle in which the compound was solubilized, and is PBS containing 5% DMSO or is PBS containing Tween (2 µl/10 ml). The animals are sacrificed on day 14 and bone growth measured by histomorphometry. Bone samples for quantitation are cleaned from adjacent tissues and fixed in 10% buffered formalin for 24-48 hours, decalcified in 14% EDTA for 1-3 weeks, processed through graded alcohols; and embedded in paraffin wax. Three to five µm sections of the calvaria are prepared, and representative sections are selected for histomorphometric assessment of the effects on bone formation and bone resorption. Sections are measured by using a camera lucida attachment to trace directly the microscopic image onto a digitizing plate. Bone changes are measured on sections cut 200 µm apart, over 4 adjacent 1x1 mm fields on both the injected and noninjected sides of the calvaria. New bone is identified by its characteristic woven structure, and osteoclasts and osteoblasts are identified by their distinctive morphology. Histomorphometry software (OsteoMeasure, Osteometrix, Inc., Atlanta) is used to process digitizer input to determine cell counts and measure areas or perimeters.

Additional In Vivo Assays

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Lead compounds can be further tested in intact animals using an *in vivo*, dosing assay. Prototypical dosing may be accomplished by subcutaneous, intraperitoneal or oral administration, and may be performed by injection, sustained release or other delivery techniques. The time period for administration of test compound may vary (for instance, 28 days as well as 35 days may be appropriate). An exemplary, *in vivo* subcutaneous dosing assay may be conducted as follows:

In a typical study, 70 three-month-old female Sprague-Dawley rats are weightmatched and divided into seven groups, with ten animals in each group. This includes a baseline control group of animals sacrificed at the initiation of the study; a control group administered vehicle only, a PBS-treated control group, and a positive control group administered a compound (nonprotein or protein) known to promote bone growth. Three dosage levels of the compound to be tested are administered to the remaining three groups.

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Briefly, test compound, positive control compound, PBS, or vehicle alone is administered subcutaneously once per day for 35 days. All animals are injected with calcein nine days and two days before sacrifice (two injections of calcein administered each designated day). Weekly body weights are determined. At the end of the 35-day cycle, the animals are weighed and bled by orbital or cardiac puncture. Serum calcium, phosphate, osteocalcin, and CBCs are determined. Both leg bones (femur and tibia) and lumbar vertebrae are removed, cleaned of adhering soft tissue, and stored in 70% ethanol for evaluation, as performed by peripheral quantitative computed tomography (pqCT; Ferretti, J. Bone (1995) 17:3538-648), dual energy X-ray absorptiometry (DEXA; Laval-Jeantet A. et al. Calcif Tissue Intl (1995) 56:14-18; J. Casez et al. Bone and Mineral (1994) 26:61-68) and/or histomorphometry. The effect of test compounds on bone remodeling can thus be evaluated.

Lead compounds also be tested in acute ovariectomized animals (prevention model) using an *in vivo* dosing assay. Such assays may also include an estrogentreated group as a control. An exemplary subcutaneous dosing assay is performed as follows:

In a typical study, 80 three-month-old female Sprague-Dawley rats are weightmatched and divided into eight groups, with ten animals in each group. This includes a baseline control group of animals sacrificed at the initiation of the study; three control groups (sham ovariectomized (sham OVX) + vehicle only; ovariectomized (OVX) + vehicle only, PBS-treated OVX); and a control OVX group that is administered a compound known to promote bone growth. Three dosage levels of the compound to be tested are administered to the remaining three groups of OVX animals.

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Since ovariectomy (OVX) induces hyperphagia, all OVX animals are pair-fed with sham OVX animals throughout the 35 day study. Briefly, test compound, positive control compound, PBS, or vehicle alone is administered subcutaneously once per day for 35 days. Alternatively, test compound can be formulated in implantable pellets that are implanted for 35 days, or may be administered orally, such as by gastric gavage. All animals, including sham OVX/vehicle and OVX/vehicle groups, are injected intraperitoneally with calcein nine days and two days before sacrifice (two injections of calcein administered each designated day, to ensure proper labeling of newly formed bone). Weekly body weights are determined. At the end of the 35-day cycle, the animals' blood and tissues are processed as described above.

Lead compounds may also be tested in chronic OVX animals (treatment model). An exemplary protocol for treatment of established bone loss in ovariectomized animals that can be used to assess efficacy of anabolic agents may be performed as follows. Briefly, 80 to 100 six month old female, Sprague-Dawley rats are subjected to sham surgery (sham OVX) or ovariectomy (OVX) at time 0, and 10 rats are sacrificed to serve as baseline controls. Body weights are recorded weekly during the experiment. After approximately 6 weeks of bone depletion (42 days), 10 sham OVX and 10 OVX rats are randomly selected for sacrifice as depletion period controls. Of the remaining animals, 10 sham OVX and 10 OVX rats are used as placebo-treated controls. The remaining OVX animals are treated with 3 to 5 doses of test drug for a period of 5 weeks (35 days). As a postitive control, a group of OVX rats can be treated with an agent such as PTH, a known anabolic agent in this model (Kimmel et al. Endocrinology (1993) 132:1577-84). To determine effects on bone formation, the following procedure can be followed. The femurs, tibiae and lumbar vertebrae 1 to 4 are excised and collected. The proximal left and right tibiae are used for pqCT measurements, cancellous bone mineral density (BMD) (gravimetric determination), and histology, while the midshaft of each tibiae is subjected to cortical BMD or histology. The femurs are prepared for pqCT scanning of the midshaft prior to biomechanical testing. With respect to lumbar vertebrae (LV), LV2 are processed

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for BMD (pqCT may also be performed); LV3 are prepared for undecalcified bone histology, and LV4 are processed for mechanical testing.

Nature of the Compounds Useful in the Invention

All of the compounds of the invention contain two aromatic systems, Ar¹ and Ar², spaced apart by a linker at a distance of 1.5-15Å, and may preferably contain at least one nitrogen atom. A summary of the structural features of the compounds included within the invention is shown in Figure 1.

As shown, Ar¹ and Ar² may include various preferred embodiments. These are selected from the group consisting of a substituted or unsubstituted aromatic ring system containing a 5-membered heterocycle; a substituted or unsubstituted aromatic ring system containing a six-membered heterocycle; a substituted or unsubstituted naphthalene moiety, and a substituted or unsubstituted benzene moiety. There are 16 possible combinations of these embodiments, if Ar¹ and Ar² are considered distinguishable. As will be clear, however, the designation of one aromatic system as Ar¹ and the other as Ar² is arbitrary; thus there are only ten possible combinations. However, for simplicity, Ar¹ and Ar² are designated separately with the realization that the choice is arbitrarily made. All linkers described herein if not palindromic, are considered to link Ar¹ to Ar² or vice-verxa whether or not the complementary orientation is explicitly shown (as it is in some cases). Thus, if Ar¹ and Ar² are different and a linker is specified as -CONR-, it is understood that also included is the linker -NRCO- when the designations Ar¹ and Ar² are retained

The noninterfering substituents on the aromatic system represented by Ar^1 and the noninterfering substituents on the aromatic system represented by Ar^2 are represented in the formulas herein by R^* and R^b , respectively. Generally, these substituents can be of wide variety. Among substituents that do not interfere with (and in some instances may be desirable for) the beneficial effect of the compounds of the invention on bone in treated subjects are included alkyl (1-6C, preferably lower alkyl 1-4C), including straight or branched-chain forms thereof, alkenyl (1-6C, preferably 1-4C), alkynyl (1-6C, preferably 1-4C), all of which can be straight or branched chains

or are aryl (6-10C) or alkylaryl (6-15C) or aryl alkyl (6-15C) and may contain further substituents. R* and R* may also include halogens, (e.g. F, Cl, Br and I), siloxy, OR, SR, NR2, OOCR, COOR, NCOR, NCOOR, and benzoyl, CF3, OCF3, SCF3, N(CF3)2. NO, NO2, CN, SO, SO2R, SO3R and the like, wherein R is alkyl (1-6C) or is H. Similarly, these substituents may contain R* as a substitute for R wherein R* is aryl (6-10C) or alkylaryl (6-15C) or aryl alkyl (6-15C). Where R* or R* substituents are in adjacent positions in the aromatic system, they may combine to form a ring. Further, rings may be included in substituents which contain sufficient carbon and heteroatoms to provide this possibility.

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The choice of noninterfering substituents depends on the overall nature of the system. For example, in compounds of the invention wherein two pyridine rings are linked through a saturated flexible linker, a CF₃ substituent para to the linker in each of the pyridine rings is particularly preferred. In those systems wherein a quinoline is coupled through a flexible conjugated or nonconjugated linker to a phenyl substituent or to a naphthyl substituent, an amino group para to the linker in the phenyl or naphthyl moiety is preferred. Particularly preferred amino groups are dimethylamino and diethylamino. In systems wherein a benzothiazole is coupled to phenyl through a flexible linker, preferred substituents on the phenyl moiety include alkoxy or alkylthio in combination with halo, in particular, chloro. Also preferred is the presence of a diethylamino group in the phenyl moiety para to the position that is coupled to the linker. In general, the presence of a substituent in the phenyl moiety para to the position of joinder to the linker is preferred.

Generally, preferred noninterfering substituents include hydrocarbyl groups of 1-6C, including saturated and unsaturated, linear or branched hydrocarbyl as well as hydrocarbyl groups containing ring systems; halo groups, alkoxy, hydroxy, amino, monoalkyl- and dialkylamino where the alkyl groups are 1-6C, CN, CF₃, OCF₃ and COOR, and the like.

Although the number of R^a and R^b may typically be 0-4 (m) or 0-5 (n) depending on the available positions in the aromatic system, preferred embodiments

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include those wherein the number of R^{a} is 0, 1 or 2 and of R^{b} is 0, 1, 2 or 3, particularly 1 or 2.

The linker group, L, may be a covalent bond or any group having a valence of at least two and covering a linear distance of from about 1.5 to about 15 Angstroms, including those that contain cyclic moieties, that meet this spatial requirement. Useful linkers are divided, by definition herein, into three general categories: (1) flexible nonconjugating linkers, (2) flexible conjugating linkers, and (3) constrained linkers.

The preferred choice of linker will depend on the choices for Ar¹ and Ar².

As defined herein, flexible nonconjugating linkers are those that link only one position of Ar1 to one position of Ar2, and provide only a single covalent bond or a single chain between Ar1 and Ar2. The chain may contain branches, but may not contain π -bonds (except in the branches) or cyclic portions in the chain. The linker atoms in the chain itself rotate freely around single covalent bonds, and thus the linker has more than two degrees of freedom. Particularly useful flexible nonconjugating linkers, besides a covalent bond, are those of the formulas: -NR-, -CR2-, -S-, or -O-, wherein R is H or alkyl (1-6C), more preferably H or lower alkyl (1-4C) and more preferably H. Also contemplated are those of the formulas: -NRCO-, -CONR--CR2S-, -SCR2-, -OCR2-, -CR2O-, -NRNR-, -CR2CR2-, -NRSO2-, -SO2NR-, -CR2CO-, -COCR2-, and -NR-NR-CO-CR2- and its complement -CR2-CO-NR-NR-. or -NRCR2CR2NR- or the thiolated counterparts, and particularly -NHCR2CR2NH-, including the isosteres thereof, such as -NRNRCSNR- and -NRNRCONR-. Also contemplated are those of the formulas: -NH(CH2)2NH-, -O(CR2)2O-, and -S(CR₂)₂S-, including the isosteres thereof. The optimum choice among flexible nonconjugating linkers is dependent on the nature of Ar1 and Ar2.

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-NR-N=CR-; -NR-NR-CO-CR=CR-, -N=NCOCR₂-, -N=NCSCR₂-, -N=NCOCR₂-CR₂, -N=NCONR-, -N=NCSNR-, and the like, where R is H or alkyl (1-6C), preferably H or lower alkyl (1-4C); and more preferably H.

Constrained linkers are those that have more than one point of attachment to either or both Ar¹ and Ar² and, thus, generally allow for only one degree of freedom. Constrained linkers most frequently form fused 5- or 6-membered cyclic moieties with Ar¹ and/or Ar² where either Ar¹ or Ar² has at least one substituent appropriately positioned to form a second covalent bond with the linker, e.g., where Ar² is a phenyl group with a reactive, ortho-positioned substituent, or is derivatized to the linker directly at the ortho position. (Although the aromatic moieties should properly be referred to as phenylene or naphthylene in such cases, generally the term "phenyl" or "naphthyl" is used herein to include both monovalent and bivalent forms of these moieties.) Examples of particularly useful constrained linkers include

15 and the like, where X is O, N, S or CR, and Y is CR2 or C=O.

In one class of preferred embodiments, Ar¹ is an aromatic system containing a 5-membered heterocycle, of the formula:

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wherein Z is S, O, NR or $-CR_2$ in formula (1a) or CR in formula (2a), where each R is independently H or alkyl (1-6C), the dotted line represents an optional π -bond, each R^* is independently a noninterfering substituent as defined above, and m is an integer of 0-4.

In general, Ar^2 is phenyl, naphthyl, or an aromatic system containing a 5- or 6-membered heterocyclic ring. All may be unsubstituted or substituted with noninterfering substituents, R^b .

When Ar² is an aromatic system containing a six-membered heterocycle, the 10 formula of said system is preferably:

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$$\begin{array}{c|c}
R^{b}_{m} & z = z \\
z = z & (iv)
\end{array}$$

wherein each Z is independently a heteroatom selected from the group consisting of S, O and N; or is CR or CR_2 , the dotted lines represent optional π -bonds, each R^k is independently a noninterfering substituent, and m is an integer of 0-4, with the proviso that at least one Z must be a heteroatom.

Ar2 in these compounds may also have the formula

where R^b is a noninterfering substituent as defined above and n is an integer from 0 to 5.

Similarly, when Ar² is naphthyl, it may contain 0-5 R^b substitutions. When Ar² is an aromatic system containing a 5-membered heterocycle, preferred forms are those as described for Ar¹.

Thus, in one set of preferred compounds. Ar1 is

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wherein each R^* is a noninterfering substituent, m is an integer of 0-4, the dotted line represents an optional π bond, and Z is O, S, NR or CR₂ in formula (1) or is CR in formula (2) wherein each R is independently H or alkyl (1-6C).

In one group of these compounds, L is a flexible conjugating or nonconjugating linker. In this group, when Z is NR, Ar² is preferably a substituted or unsubstituted aromatic system containing a 5-membered heterocycle or is

wherein R^b is a noninterfering substituent and n is an integer of 0-5; and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR-where R is H or alkyl (1-6C); and/or the dotted line represents a π bond.

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In these embodiments as well as in alternative embodiments of Ar^2 , it is preferred that each R^b is independently halo, OR, SR, NR2, NO, NO2, OCF3 or CF3 wherein R is H or alkyl (1-6C), or R^b comprises an aromatic system.

Preferred compounds in this group are 59-0100, 59-103, 59-104, 59-105 and 59-106 (See Figure 13).

In another group of these compounds with flexible linkers, Z is S, and AT^2 is preferably a substituted or unsubstituted aromatic system containing a 6-membered heterocycle or is of the formula

wherein R^b is a noninterfering substituent and n is an integer of 0-5; and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR-where R is H or alkyl (1-6C); and/or the dotted line represents a π bond.

In such compounds, regardless of the choice of Ar², preferred are those compounds wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.

Both when Z is S and when Z is NR, it is preferred that m is 0 and/or each R^b is independently OR, SR or halo, where n=2 and at least one R^b is independently OR or SR and/or L is -NHCO- or -CR=-CR-

Preferred compounds in this group include compounds 59-002, 59-0070, 59-0072, 59-0099, 59-0102, the benzothiazole counterpart of 59-0104, 59-0144, 59-0147, 59-0149, 59-0186, 59-0187, 59-0192, 59-0193, 59-0195, 59-0197, 59-0202, 59-0204, 59-0205, 59-0206, 59-0207, 59-0208, and 59-0210, especially the benzothiazole counterpart of 59-0104 or compounds 59-0147, 59-0205 or 59-0210. (See Figure 13)

Z can also be CR, CR2 or O; here it is also preferred that Ar2 is

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wherein R^b is a noninterfering substituent and n is an integer of 0-5, and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR-where R is H or alkyl (1-6C), and/or the dotted line represents a π bond.

In these compounds, too, it is preferred that each R^b is independently halo, OR, SR, NR_2 , NO, NO_2 , OCF_3 or CF_3 wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system. A preferred compound is 896-5005. (See Figure 4)

The compounds wherein Ar¹ is 1a or 2a as above may also contain a constrained linker.

In these compounds, preferred Z is S or NR; and/or those wherein L is selected from the group consisting of

Ar2 is

wherein Rb is a noninterfering substituent and m is 0-4.

Preferably, each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system. A preferred compound is 59-0124. (See Figure 13)

In another group of preferred embodiments, Ar1 is of the formula

wherein each R^a is independently a noninterfering substituent or is H and Z is NR, S or O, wherein R is alkyl (1-6C) or H, especially where Z is S and/or wherein Ar^2 is

wherein R^b is a noninterfering substituent and n is an integer of 0-5., and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR-where R is H or alkyl (1-6C), and/or the dotted line represents a π bond. Especially preferred are those compounds where each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.

In another group of compounds, Ar1 is

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$$R_{m}^{a}$$
 (4a)

wherein R^* is a noninterfering substituent, m is an integer of 0-4, each dotted line represents an optional π -bond, each Z is independently N, NR, CR or CR_{2*} where each R is independently H or alkyl (1-6C) with the proviso that at least one Z is N or NR.

Particularly preferred members of this group are those wherein Ar1 is

especially those wherein Ar2 is

$$R^b_n$$
 R^b_m R^b_m (vi) or N (via)

wherein each R* is independently a noninterfering substituent, and n is 0-5 and m is 0-4, and/or L is -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR2-, -NRCR2CR2-, -NRCR2CO-, -NRNR-, -CR2CR2-, -NRCR2CR2NR-, -NRCR=CRNR- or -NRCOCR3NR-.

In general, preferably each R^b is independently halo, OR, SR, NR₂, NO, NO₂,
OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.

In an especially preferred group, m is 0, each R^b is NR₂ or OR and n is 1 or 2,
and/or L is -CR=CR-, -N=N- or -NRCO-, especially the compounds of formulas
59-0030, 59-0078, 59-0091, 59-0093, 59-0150, 50-0197, 59-0198, 59-0199 or
59-0480. (See Figure 13)

Also preferred are those wherein ${\rm Ar}^1$ has formula (4a) or (5a) and wherein ${\rm Ar}_2$ is substituted or unsubstituted quinolyl or naphthyl of the formula

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wherein each R^b is a noninterfering substituent and m is 0-4.

Preferred among these are those wherein L is -N=N-, -RC=CR-, -RC=N-,
-NRCO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CO-, -NRNR-, -CR₂CR₂-,
-NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-, and/or wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system and m is 0, 1 or 2.

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The compounds 59-0089, 59-0090, 59-0092 or 59-0094 are particularly preferred.

Ar1 is also preferably

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5 wherein each R* is a noninterfering substituent and m is 0-4, in particular where L is -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CO-, -NRNR-, -CR₂CR₂-, -NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-, and/or Ar² is

wherein R^b is a noninterfering substituent and n is an integer of 0-5. Especially preferred are compounds wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system, in particular compounds 59-203, 59-285 or 59-286. (See Figure 13)

When Ar¹ is of formula (4a), L can also be a constrained linker. In still another preferred set, Ar¹ is

$$\begin{array}{c|c}
R^{a}_{m} & z = z \\
z & z \\
z - z
\end{array} (9a)$$

wherein each R* is independently a noninterfering substituent, m is an integer of 0-4, each Z is independently N or CR, where R is H or alkyl (1-6C), with the proviso that at least one Z must be N and at least one Z must be CR.

In these compounds, L is preferably a flexible conjugating or nonconjugating linker, and/or wherein Ar² is

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$$R_{m}^{b}$$
 (v) or $Z = Z$ (vi)

wherein each \mathbb{R}^b is independently a noninterfering substituent, and in (vi) each Z is independently \mathbb{N} or $\mathbb{C}\mathbb{R}$, where \mathbb{R} is \mathbb{H} or alkyl (1-6C), with the proviso that at least one \mathbb{Z} must be a \mathbb{N} and at least one \mathbb{Z} must be $\mathbb{C}\mathbb{R}$.

Preferred such compounds have the formula

$$R_{m}^{a}$$
 or R_{m}^{a} L R_{m}^{b} or R_{m}^{a}

Preferred L embodiments in this group include -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR₂-, -NRCR₂CC₂-, -NRCR₂CO-, -NRNR-, -CR₂CR₂-, -NRCR₂CR₂-, -NRCR₂CO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-; preferred for R* and R* are halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R* or R* comprise aromatic systems and each m and n is independently 0. 1 or 2.

In particular, compounds are preferred where L is -NHCR₂CR₂NH- and R^{*} is CF₃ para to L, especially compounds 59-0145, 59-0450, 59-0459 or 59-0483. (See Figure 13)

Finally, in another preferred group, Ar1 is

wherein each R³ is a noninterfering substituent, and n is an integer of 0 and 5, and wherein L is a flexible linker that contains at least one nitrogen. In the alternative or in addition. Ar² is of the formula

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and L is -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR₂-, -NRCR₂CR₂-,
-NRCR₂CO-, -NRNRCR₂CR₂-, -NRNRCB=CR-, -NRNRCOCR₂-,
-NRNRCOCR=CR-, -NRNRCSCR₂-, -NRNRCSCR=CR-, -NRNRCONR-,
-NRNRCSNR-, -NRNR-, -CR₂CR₂-, -NRCR₂CR₂NR-, -NRCR=CRNR- or
-NRCOCR₂NR-. It is preferred that each R⁸ is independently halo, OR, SR, NR₂, NO,
NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R⁸ comprises an aromatic system.

Especially preferred are those compounds wherein L is -CR=CRCONRNR-,
-CR=CRCSNRNR-, -CR₂CONRNR- -CR₂CSNRNR-, -NRNRCONR- or
-NRNRCSNR- and/or R⁸ is -NR₂ and n=1 wherein R⁸ is in the para position, especially wherein R⁸ is -COOR and m is 1; most especially compounds 59-0045, 59-0095,
59-0096, 59-0097 and 59-0098. (See Figure 13)

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As set forth above, several families of preferred embodiments are defined by specifying Ar^1 and Ar^2 , and L. In one such family, wherein Ar^1 is an aromatic system containing a 5-membered heterocyclic ring, the compound 59-0072, wherein Ar^1 is unsubstituted benzothiazole, the linker $(Ar^1 \rightarrow Ar^2)$ is NHCO, and Ar^2 is 2-methoxy-4-methylthiophenyl was used as a lead compound and variations of the structure studied. Figure 5 shows representative compounds synthesized to analyze the effects of the nature of the linker, various alternatives of Ar^1 wherein Z is O, NR or S, and the effect of substitution on the phenyl moiety, as well as the heterocycle.

Figure 5 gives the structures of these compounds, along with their maximum activity as compared to 59-0008 at 10 μM (the maximum for 59-0008) in the *in vitro* bone growth stimulation assay as well as the concentration at which 50% of maximum stimulation of the BMP promoter was obtained (EC₅₀). See Example 1 for the details of this assay. The results of this study indicate that the amide linker in 59-0072 can readily be substituted by -CH=CH- and that the substitution on the phenyl ring had advantageous effects in the order: 2-Cl-4-OMe=2,4-di-OMe=2-OMe-4-SMe >>3,4-di-OMe=4-OMe. In general, compounds 59-0205, 59-0104, 59-0107, 59-0210 and 59-0124 have the best activity in the primary screen, but only 59-0124 is active in the *ex vivo* calvarial assay described in Example 3.

Similar structure/activity relationship studies were conducted for compounds wherein Ar¹ is quinoline. In this study, compound 50-0197, wherein Ar¹ is unsubstituted quinoline, the linker is -CH=CH-, and Ar² is p-dimethylaminophenyl was used as a lead compound. The compounds synthesized in this study are shown in Figure 6, along with their maximum stimulation characteristics and EC₅₀ in the assay of Example 1. The results of these studies showed that quinoxaline analogs are the most active in the assay, followed by quinoline; the linker can most preferably be -CH=CH-or -N=N- as judged by activity in the assay, but -CH=CH- is preferred in vivo due to its lack of toxicity. Preferred substituents on the phenyl ring in Ar² include 2,4-di-OMe; 4-NMe₂-2-OMe, and 4-NMe₂. For the compounds in Figure 6, 59-0282 and 50-0197 were moderately active and 59-0203 was highly active in the ex vivo calvarial assay described hereinabove as a modification of Gowen, M. and Mundy, G. J. Immunol (1986) 136:2478-2482.

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Another group of compounds wherein Ar¹ and Ar² are pyridyl heterocycles was also studied. In this case, compound 59-0145 was used as the lead compound; the linker, the nature of the substituents R⁴ and R^b were varied. In one instance, a quinolyl residue was substituted for a pyrimidine residue as Ar². Representative compounds used in this study are shown in Figure 7, along with the data from the screening assay.

Using 59-0145 as a lead, a CF_3 group in one of Ar^1 and Ar^2 appeared essential; however, one of R^4 or R^5 could also be NO_2 or CN. The most preferred linker is -NHCH₂CH₂NH-; substitution on the amino groups in L by an alkyl group appeared to reduce activity. Enhanced chain lengths also led to loss of activity.

Preferred compounds in this group, which perform better than 59-0008 in the screening assay, included 59-0450, 59-0459, 59-0480, and 59-0483.

Finally, a series in which Ar¹ is 3-carboxyphenyl was studied using 59-0045 as the lead compound. In 59-0045, L is -NHN=CH- and Ar² is p-dimethylaminophenyl. Figure 8 shows the compounds synthesized in this series. Under the circumstances of this assay, analogs wherein R^b was, instead of a nitrogen-containing moiety, F, Cl, or OMe were inactive. Preferred compounds in this series are 59-0096 and 59-0098. 59-0098 is very active in the ex vivo calvarial assay described above.

Synthesis of the Compounds Useful in the Invention

Many of the compounds useful in the invention are commercially available and can be synthesized by art-known methods. Those compounds useful in the invention which are new compounds, can similarly be obtained by methods generally known in the art, as described in the Examples below.

The following examples are intended to illustrate, but not to limit, the invention.

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Preparation A

Compound 59-0008 used as a standard in the assays, was synthesized according to the procedure of McDonald, W. S., et al. Chem Comm (1969) 392-393: Irving, H. N. N. H. et al. Anal Chim Acta (1970) 49:261-266. Briefly, 10.0 g of 15 dithizone was taken up in 100 ml EtOH and 50 ml AcOH and heated at reflux for 18 h. After cooling, this was diluted first with 100 ml water and then with 50 ml 1N NaOH. This was then further neutralized by the addition of 6 N NaOH to bring the pH to 5.0. This deep purple mixture was then concentrated on a rotavapor to remove organics. Once the liquid had lost all of its purple color, this was filtered to collect the dark precipitate. Purification by flash chromatography (4.5 x 25.7 cm; EtAc/Hep. (1 4); Rf 20 0.22) followed by recrystalization from EtOH gave 2.15 g (25% yield) of dark purple crystals, mp=184-185 °C. 1H NMR (CDCl3) 7.90 (d of d, J1=7.7, J2=2.2, 2H), 7.64 (hump, 1H), 7.49 (m, 3H), 7.02 (m, 1H), 6.91 (m, 2H), 6.55 (d, J=8.1, 1H). MS (EI) 254 (47, M+), 105 (26), 77 [100], 51 (27). HRMS (EI, M+) 254.0626 (calcd 25 254.0626182). Anal. Calcd for C13H10N4S: C, 61.40; H, 3.96; N, 22.03. Found: C, 61.40; H, 4.20; N, 22.06.

Example 1

High Throughput Screening

Several tens of thousands of compounds were tested in the assay system set forth in WO 96/38590, published 5 December 1996, and incorporated herein by reference. The standard positive control was 59-0008 (also denoted "OS8"), which is of the formula:

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In more detail, the 2T3-BMP-2-LUC cells, a stably transformed osteoblast cell line described in Ghosh-Choudhury et al. Endocrinology (1996) 137:331-39. referenced above, was employed. The cells were cultured using α-MEM, 10% FCS with 1% penicillin/streptomycin and 1% glutamine ("plating medium"), and were split 1:5 once per week. For the assay, the cells were resuspended in a plating medium containing 4% FCS, plated in microtiter plates at a concentration of 5 x 103 cells (in 50 μl)/well, and incubated for 24 hours at 37°C in 5% CO2. To initiate the assay, 50 μl of the test compound or the control in DMSO was added at 2X concentration to each well, so that the final volume was 100 μl. The final serum concentration was 2% FCS. and the final DMSO concentration was 1%. Compound 59-0008 (10 µM) was used as a positive control.

The treated cells were incubated for 24 hours at 37°C and 5% CO2. The medium was then removed, and the cells were rinsed three times with PBS. After 20 removal of excess PBS, 25 µl of 1X cell culture lysing reagent (Promega #E153A) was added to each well and incubated for at least ten minutes. Optionally, the plates/samples could be frozen at this point. To each well was added 50 ul of luciferase substrate (Promega #E152A; 10 ml Promega luciferase assay buffer per 7 mg Promega luciferase assay substrate). Luminescence was measured on an

automated 96-well luminometer, and was expressed as either picograms of luciferase activity per well or as picograms of luciferase activity per microgram of protein.

In this assay, compound 59-0008 (3-phenylazo-1H-4,1,2-benzothiadiazine) exhibited a pattern of reactivity, as shown in Figure 2. The activity for compound 59-0008 was maximal at a concentration of approximately 3-10 μ M and, more particularly, at about 3 μ M, and thus provided a response of approximately 175 light emission units. Accordingly, other tested compounds were evaluated at various concentrations, and these results were compared to the results obtained for 59-0008 at 10 μ M (which value was normalized to 100). For instance, any tested compound in Figure 3 and Figure 4 that showed greater activity than 10 μ M of 59-0008 would result in a value over 100.

As shown in Figure 3 (46 sheets) and Figure 4 (28 sheets), several compounds were found to be particularly effective.

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Example 2

In vivo Calvarial Bone Growth Data

Compound 59-0008 was assayed in vivo according to the procedure described previously (see "In vivo Assay of Effects of Compounds on Murine Calvarial Bone Growth", supra). As compared to a vehicle control, compound 59-0008 induced a 4-fold increase in width of new calvarial bone.

In another experiment, 5 week old Swiss white mice were injected 3 times a day for 5 days over the calvaria with compound 59-0203 using PBS, 5% DMSO and 0.1% BSA as carrier. The drug was tested at 6 different doses, from 0.1-50 mg/kg/day. Animals were sacrificed 3 weeks after the injections started and calvariae were fixed, decalcified, and processed for histology. Bone histomorphometry measuring total bone area (BA/TV) confirms that FGF, used in every experiment as a positive control, shows an increase in the total bone area with all doses tested, but this increase is only significantly different from control at 1 and 5 mg/kg/day. The invention compound 59-0203 shows consistent increases over the 0.1-50 mg/kg/day range at a somewhat lower level than that obtained with FGF.

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Similar results are obtained when new bone width in microns is measured. There was no new bone present in the control group. 59-0203 caused new bone formation at all doses, with a significant increase at 25-50 mg/kg/day. New bone as percentage of the total bone area was about 45% for the FGF positive control and from about 15% to 30% over the range of 0.1-50 mg/kg/day for 59-0203. There was no new bone present in the negative control.

Example 3

Ex vivo Calvarial Bone Growth Assay

A number of compounds, in particular, those studied in connection with lead compounds classified as hydrazone/hydrazides (H) exemplified by 59-0045, benzothiazoles (T) exemplified by 59-0104, bis-pyridines (P) exemplified by 59-0145, and quinolines/quinoxalines (Q) exemplified by 59-0197, were tested in the ex vivo calvarial assay described hereinabove. The results of this assay are shown in Figure 9. In this assay, histomorphotometry and osteoblast numbers are measured and effects are measured on an arbitrary scale from 1-3: i.e., 1, 1+, 2-, 2, 2+, 3-, 3, wherein 1 denotes "inactive." In this assay, for example, FGF scores 2-3.

The scores are assigned to bone formation on the ectocranial periosteal surface.

The area immediately surrounding midline suture is excluded from analysis.

Score

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- 0 Toxicity. Cell necrosis, pyknotic nuclei, matrix disintegration.
- 1 A score of "1" is the bone forming activity seen in control cultures containing BGJb media +0.1% bovine serum albumin. The periosteal surface is covered by one layer of osteoblasts (at about 50% of the bone surface, with the remaining 50% being covered by bone lining cells). A score of "1-" is assigned if less than 50% of the periosteal surface is covered by osteoblasts due to inhibitory activity or minor toxicity of the agents being tested. A score of "1+" is given if over 50% of the surface is covered by osteoblasts.
- 2 A moderate increase in bone forming activity. 20-40% of the periosteal surface is covered by up to two layers of osteoblasts. A score of "2-" is given if less than 20% of the surface is covered by

two layers and "2+" if more than 40% of the surface is covered by two layers of osteoblasts.

3 A score of "3" is the bone forming activity seen in control cultures containing BGJb media + 0.1% BSA +10% fetal bovine serum. More than 20% of the periosteal surface is covered by three layers of osteoblasts. The cells appear plump (size can exceed 100µm2). A score of "3-" is given if less than 20% of the periosteal surface is covered by three layers of osteoblasts and or osteoblast size is less than 100µm2. A score of "3-" has never been observed.

In all samples, toxicity, ectopic new or woven bone formation associated with osteoblasts, and osteoblast size as reflections of relative activity are noted.

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The results shown in Figure 9 represent those obtained when the measurements were made by two different groups. It is clear that a number of compounds tested have activity in this assay. From the results shown in Figure 9, 59-0073, 59-0030, 59-0070, 59-007, 59-0019, 59-0099, 59-0072 and 59-0103 show at least some indication of activity. 59-150 and 59-0104 showed activity when measured by one group but not the other, similarly, 50-0197 had this pattern. It appears that 59-0098 and 59-0203 are quite active in this assay and 59-0145 shows a consistent moderate activity.

Example 4

Stimulation of Bone Growth in Ovariectomized Rats (OVX Assay)

The compound 59-0145 was tested at various concentrations in the OVX assay conducted as described above. The increase in bone volume was measured by two different groups; one group found 5 µg/kg/day of 59-0145 gave 21% increase over control whereas the second group found a 71% increase. At 50 µg/kg/day, the first group found a 31% increase, and the second a 54% increase.

In another experiment, the lumbar vertebrae were measured and the above dosages of 59-0145 were shown to provide a beneficial effect, as shown in Figure 10.

In another experiment, 3 month old Sprague Dawley rats were ovariectomized and depleted for six weeks. At the end of the six weeks, treatment was started with subcutaneous administration of compound 59-0145. The treatment continued for 10

weeks. At the end of the 10 weeks animals were sacrificed, bones were collected for qCT measurements and histology; serum was also collected for osteocalcin determinations

Figure 11 shows the percentage increase in trabecular bone (proximal tibia) compared to the placebo-treated group in chronic ovariectomized rats after 10 weeks of treatment. Compound 59-0145 causes significant increase in trabecular bone at doses of 50-500 wa/ka/day.

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Figure 12 shows results of qCT and bone histomorphometry in proximal tibia in the first two panels, as well as serum osteocalcin levels at the time of sacrifice as a percentage increase compared to control group (OVX placebo-treated group).

Example 5

Chondrogenic Activity

Compounds 59-008, 59-0102 and 50-0197 were assayed for effects on the differentiation of cartilage cells, as compared to the action of recombinant human BMP-2. Briefly, a mouse clonal chondrogenic cell line, TMC-23, was isolated and cloned from costal cartilage of transgenic mice containing the BMP-2 gene control region driving SV-40 large T-antigen, generated as described in Ghosh-Choudhury et al Endocrinology 137:331-39, 1996. These cells were cultured in DMEM/10% FCS, and were shown to express T-antigen, and also to produce aggrecan (toluidine blue staining at pH 1.0) and Type-II collagen (immunostaining) by 7 days after confluence.

For measurement of alkaline phosphatase (ALP) activity, the technique of LF Bonewald et al. J Biol Chem (1992) 267:8943-49, was employed. Briefly, TMC-23 cells were plated in 96 well microtiter plates in DMEM containing 10% FCS at 4 x 10³ cells/well. Two days after plating, the cells were confluent and the medium was replaced with fresh medium containing 10% FCS and different concentrations of compounds or recombinant BMP-2. After an additional 2 or 5 days incubation, the plates were washed twice with PBS, and then lysing solution (0.05% Triton X-100) was added (100 µl/well). The cells were lysed by three freeze-thaw cycles of -70°C (30 min), followed by 37°C (30 min with shaking). Twenty microfiters of cell lysates

were assayed with 80 μ l of 5 mM p-nitrophenol phosphate in 1.5 M 2-amino-2-methyl-propanol buffer, pH 10.3 (Sigma ALP kit, Sigma Chemical Co., St. Louis, MO) for 10 min at 37°C. The reaction was stopped by the addition of 100 μ l of 0.5 M NaOH. The spectrophotometric absorbance at 405 nm was compared to that of p-nitrophenol standards to estimate ALP activity in the samples. The protein content of the cell lysates was determined by the Bio-Rad protein assay kit (Bio-Rad, Hercules, CA). Specific activity was calculated using these two parameters.

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product was collected by filtration.

At day 2, compounds 59-0008 (10° M), 59-0102 (10° M) and 59-0197 (10° M) increased ALP levels approximately 3-, 2- and 2.5-fold, respectively, as compared to the vehicle control. Recombinant BMP2 at 100, 50 or 10 ng/ml induced ALP levels approximately 10-, 4- or 1.5-fold, respectively, as compared to the vehicle control.

Example 6

Synthesis of Exemplary Compounds

- 15 A. Compounds of the invention wherein Ar¹ is of formula (1a) or (2a) can be synthesized by the procedures described in Dryanska, V. and Ivanov, K. Synthesis (1976) 1:37-8, using the described embodiments of Ar² and the appropriate analogous heterocycle embodied in Ar¹ substituted for the benzothiazole shown. Alternates to the olefin linker described can also be prepared using standard methods.
- 20 Compounds of the invention represented by exemplary Compound 59-0234, wherein Z is O, L is -CH=CH-, and Ar² is 2,4-dimethyoxy-phenyl, including Compounds 59-0211 and 59-0233, were prepared according to the following procedure describing synthesis of Compound 59-0234. Briefly, to a N,N-dimethylformamide (DMF) solution of 2-methylbenzoxazole (1 mmol) and 25 2,4-dimethoxybenzaldehyde (1 mmol) was added lithium t-butoxide (2 mmol). The reaction mixture was heated at 130°C for 3h. After cooling to room temperature, the reaction mix was poured into ether and washed several times with water. The organic phase was dried over Na₂SO₄, filtered. and evaporated to dryness. The residue was dissolved in a minimal amount of hot ether and, on standing overnight, the crystalline

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PCT/US97/18864

B. Exemplary Compound 59-0150 where Ar¹ is of formula 4a was synthesized according to the procedure of Zamboni et al. J Med Chem (1992) 35:3832-44. First, 2-triphenylphosphoniumquinaldine bromide was synthesized as follows. Quinaldine (200 mmols), NBS (200 mmols) and a catalytic amount of benzoyl peroxide (10 mmols) were dissolved in 1 L of anhydrous carbon tetrachloride, and the mixture was stirred under reflux for 72 h. The mixture was cooled to RT and washed with water. The organic layer was drawn off, dried over anhydrous sodium sulfate, filtered and concentrated in vacuo to a dark oil. The crude mixture was dissolved in 500 ml of acetonitrile, then triphenylphosphine (200 mmols) was added and the mixture was refluxed under nitrogen overnight. It was then cooled to RT and diluted with anhydrous ether. The precipitated solid was collected by filtration, washed thoroughly with anhydrous ether and dried in vacuo overnight, yielding 25 g of a tan crystalline solid which showed a single spot by TLC (silica gel, 5 % MeOH in DCM).

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A Wittig reaction was then performed. Briefly, under anhydrous conditions, 0.738 g (1.68 mmol) 2-triphenylphosphoniumquinaldine bromide in dry THF was cooled to -78°C. 1.0 ml (2.5 mmol, 2.5 M in hexanes) n-butyl lithium was slowly added, and this was allowed to react for 20 min. 0.301 g (1.68 mmol) 4-(N,N-dimethylamino)-2-methoxybenzaldehyde was then added. After a few minutes, the cold bath was removed, and this was left at ambient temp. for 18 h. The reaction was quenched by the addition of aq. sat. NH4Cl. This was extracted with EtAc, and the organics washed with additional NH4Cl, sat. NaHCO3, and sat. NaCl. This was dried over anhydrous Na₂SO₄ and the solvent stripped on a rotavapor. After flash chromatography (3.8 x 18.0 cm, EtAc/Hep. (1.3); Rf 0.29), 0.135 g (26% yield) of a red solid was obtained, mp=185-187 °C. ¹H NMR (CDCl3) 8.04 (t, J=9.0, 2H), 7.94 (d, J=16.5, 1H), 7.74 (d, J=8.1, 1H), 7.73 (d, J=8.5, 1H), 7.66 (t of d, J₁=7.6, J_d=1.4, 1H), 7.51 (d, J=8.8, 1H), 7.43 (t of d, J₁=7.6, J_d=1.1, 1H), 7.29 (d, J=16.6, 1H), 6.37 (d of d, J₁=8.7, J₂=2.4, 1H), 6.22 (d, J=2.4, 1H), 3.93 (s, 3H), 3.03 (s, 6H). Anal. Calcd for C₂OH₂ON₂O₂C. 7.892: H, 6.62: N, 9.20. Found:

Exemplary Compound 59-0209 was synthesized according to the procedure of McOmie, J. F. W.; and West, D. E., Org Synth, Collect Vol V (1973) 412. Under anhydrous conditions, 0.510 g (1.95 mmol) NNC 59-0198 was slowly treated with 0.38 ml (3.9 mmol) BBr3 in dry CH2Cl2 at -78°C. After 15 min, this was allowed to warm to RT. After 2 h, the reaction was re-cooled to -78°C, and was then quenched by the addition of 1.6 ml (12 mmol) TEA in 25 ml MeOH. After 10 min. this was again allowed to warm to ambient temperature. After 1 h, this was concentrated to dryness on a rotavapor, and twice slurred in MeOH and re-stripped Purification by flash chromatography (3.0 x 25.6 cm; EtAc/Hep. (1:2); Rf 0.25) gave 0.20 g (41% yield) of a slightly vellow solid, mp=271-272 °C (dec.). 1H NMR

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- (DMSO-d6) 9.77 (s, 1H), 8.31 (d, J=8.6, 1H), 7.96 (d, J=8.6, 1H), 7.92 (d, J=8.3, 1H), 7.82 (d, J=8.6, 1H), 7.74 (d, J=16.6, 1H), 7.72 (t, J=7.6, 1H), 7.58 (d, J=8.6, 2H), 7.53 (t, J=7.6, 1H), 7.26 (d, J=16.5, 1H), 6.83 (d, J=8.6, 2H). Anal. Calcd for C17H13NO: C, 82.57; H, 5.30; N, 5.66. Found:
- Exemplary Compound 59-0019 was synthesized as follows: to a xylene solution of 2-methylquinoxaline (10 mmol) and 4-dimethylaminobenzaldehyde (10 mmol) was added piperdine (2 ml). The solution was heated at reflux for 1 day, at which time DBU (200 µL) was added and reflux continued for another 2 days. The solution was cooled to RT and extracted with 1 M citric acid. The aqueous phase was 20 repeatedly extracted with ether. The organic phases were pooled, dried over Na2SO4. filtered and evaporated to dryness. The residue was chromatographed on silica gel The product was eluted using 8:1:1 dicholormethane:ether: hexane. Fractions containing pure product were pooled and evaporated to dryness. The residue was triturated with ether and filtered to give the desired compound.
- 25 Exemplary Compound 59-0183 and related Compound 59-0182 were synthesized according to the following procedure. Briefly, quinaldic acid (0.5 mmol) and HATU (0.5 mmol) were dissolved in 2.5 mL of anhydrous DMF in a vial and the solution was stirred at room temperature (RT). Diisopropylethyamine (1 mmol) was added dropwise to the above stirred solution and the mixture was stirred for 15 min. 30 The appropriate amine (0.5 mmol) was then added all at once to the above stirred

mixture, and the mixture was stirred overnight at RT. It was then diluted with 25 mL of cold water with vigorous stirring, the precipitate was collected by filtration and washed thoroughly with water several times, and then dried in vacuo overnight. The product was purified by flash column chromatography over silica gel eluting with dichloromethane. The pure product was obtained as a tan powder.

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- F. Exemplary Compound 59-0209 was synthesized according to the following procedure. Under anhydrous conditions, 0.510 g (1.95 mmol) NNC 59-0198 was slowly treated with 0.38 ml (3.9 mmol) BBr3 in dry CH2Cl2 at -78°C. After 15 min, this was allowed to warm to RT. After 2 h, the reaction was re-cooled to -
- 78°C, and was then quenched by the addition of 1.6 ml (12 mmol) TEA in 25 ml MeOH. After 10 min, this was again allowed to warm to ambient temperature. After 1 h, this was concentrated to dryness on a rotavapor, and twice slurred in MeOH and re-stripped. Purification by flash chromatography (3.0 x 25.6 cm; EtAc/Hep. (1:2); Rf 0.25) gave 0.20 g (41% yield) of a slightly yellow solid, mp=271-272 °C (dec.). ¹H
- 15 NMR (DMSO-d6) 9.77 (s, 1H), 8.31 (d, J=8.6, 1H), 7.96 (d, J=8.6, 1H), 7.92 (d, J=8.3, 1H), 7.82 (d, J=8.6, 1H), 7.74 (d, J=16.6, 1H), 7.72 (t, J=7.6, 1H), 7.58 (d, J=8.6, 2H), 7.53 (t, J=7.6, 1H), 7.26 (d, J=16.5, 1H), 6.83 (d, J=8.6, 2H). Anal. Calcd for C17H13NO: C, 82.57; H, 5.30; N, 5.66. Found:
- G. Other embodiments wherein AR¹ is of formula (4a) can be synthesized
 as follows:

 - Azo derivatives may be obtained by reaction of 2-aminoquinolines with aldehydes, Morimoto, T., et al., Chem Pharm Bull (1977) 25:1607-09; Renault, J., et al., Hebd Seances Acad Sci, Ser C (1975) 280:1041-43; and Lugovkin, B. P.; Zh Obshch Khim (1972) 42:966-69.
 - c. Imino derivatives may be obtained by reaction of 2formylquinolines with anilines, Tran Quoc Son, et al., (1983) 21:22-26; Hagen,

V. et al. Pharmazie (1983) 38:437-39; and Gershuns, A. L., et al., Tr Kom Anal Khim, Akad Nauk SSSR (1969) 17:242-50.

- d. Alternatively conjugated linkers can be formed by bromination of the olefin of 50-0197 with Br₂ in AcOH followed by elimination with DBU as set forth in Zamboni et al. J Med Chem (1992) 35:3832-44.
- H. Analogs having the constrained linker depicted below:

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may be synthesized by reference to the methods described in Gorbulenko, N.V.

10 et al. Dokl Akad Nauk Ukr SSR (1991) 5:117-23, substituting the 6-membered heterocycle for benzothiazole.

Related, compounds having the constrained linker depicted below:

R= alkyl, OH

- may be synthesized by reference to the methods described in the following publications: Chaurasia, M.R. & Sharma, A.J. Acta Cienc Indica Chem (1992) 18:419-22; Kandeel, Maymona M., in Phosphorus, Sulfur, Silicon, Relat Elem (1990) 48:149-55; Salem, M.A. & Soliman, E.A. Egypt J Chem (1985) 27:779-87, Garin, J. et al. Synthesis (1984) 6:520-22, and Ayyangar N. R. et al. Dyes and Pigments (1990) 13:301-10
 - I. Exemplary Compound 59-0145 can be synthesized according to the following method. Briefly, a mixture of 2-chloro-5-trifluoromethylpyridine (15 mmol), ethylenediamine (6 mmol), and diisopropylethylamine (18 mmol) was heated at reflux for 18 h. After cooling to room temperature, the solid mass was triturated with

dichloromethane. The product was filtered and then suspended in hot EtOAc:CHCl3 (50:50, 800 mL) and filtered to remove insoluble material. The volume was reduced to ~200 mL by heating on a steam bath. On standing, crystals of pure product were deposited.

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- Related compounds may be synthesized by reference to the method described for Compound 59-0145, and by reference to the methods described in the following publications: Tzikas, A.& Carisch, C., US Patent No. 5,393,306, issued February 28 1995; Herzig, P.& Andreoli, A., EP 580554, published January 26, 1994; Pohlke, R. & Fischer, W., DE 3938561, published May 23, 1991. Analogs containing the structure O-(CH₂)_n-O may be synthesized by reference to the previous citations, as well as the following publications: Kawato, T. & Newkome, G. Heterocycles (1990) 31:1097-104: Kameko, C. & Momose, Y. Synthesis (1982) 6:465-66; Tomlin, C.D.S. et al., GB 1161492, published August 13, 1969.
- I Exemplary Compound 59-0097 and exemplary Compound 59-0201 15 were synthesized according to the following general procedure. Briefly, the isothiocyanate or isocyanate (1 mmol) was dissolved in 5 mL of anhydrous DMF in a vial and the solution was stirred at room temperature (RT). Diisopropylethyamine (2 mmol) was added dropwise to the above stirred solution followed by 3hydrazinobenzoic acid (1 mmol), and the mixture was stirred overnight at RT. It was 20 then diluted with 50 mL of cold water with vigorous stirring. The precipitate was collected by filtration, washed thoroughly with water several times, and then dried in vacuo overnight. The product was purified by flash column chromatography over silica gel eluting with 5 % methanol in dichloromethane. The pure product was obtained as a red to purple powder. The compounds of the invention are produced by substituting for at least one phenyl group the appropriate heterocycle.
 - Compounds of the class represented by exemplary Compound 59-0045 can be synthesized using standard procedures for the synthesis of phenyl hydrazones of aromatic aldehydes, as described in any organic textbook. The synthesis of exemplary Compound 59-0045 may be performed as follows. Briefly, a suspension of 3hydrazinobenzoic acid (1 mmol), p-dimethylaminobenzaldehyde (1 mmol), and AcOH

(50 μL) in EtOH:H₂O (4 mL:1 mL) was heated at 105°C in a sealed vial for 3 h. After cooling, a bright yellow solid was removed by filtration. The solid was washed with cold MeOH and then with ether to give pure product.

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- L. Exemplary Compound 59-0096 and related, exemplary Compounds 59-0098, 59-0095, 59-0107, 59-0108, 59-0109, 59-0100 and 59-0200 may be synthesized according to the following general procedure. Briefly, the appropriate carboxylic acid (1 mmol) and HATU ([O-(7-azabenzotriazol-1-yl)-1,1,3,3-tritetramethyluronium hexafluorophosphate]; 1 mmol) were dissolved in 5 mL of anhydrous DMF in a vial and the solution was stirred at room temperature (RT). Diisopropylethyamine (3 mmol) was added dropwise to the above stirred solution and the mixture was stirred for 15 min. 3-Hydrazinobenzoic acid (1 mmol) was then added all at once to the above stirred mixture and the mixture was stirred overnight at RT. It was then diluted with 50 mL of cold water with vigorous stirring and the precipitate was collected by filtration and washed thoroughly with water several times and then dried in vacuo overnight. The product was purified by flash column chromatography over silica gel eluting with 5 10 % methanol in dichloromethane. The pure product was obtained as a tan crystalline solid.
- M. Exemplary Compound 59-0097 and exemplary Compound 59-0201 were synthesized according to the following general procedure. Briefly, the isothiocyanate or isocyanate (1 mmol) was dissolved in 5 mL of anhydrous DMF in a vial and the solution was stirred at room temperature (RT). Diisopropylethyamine (2 mmol) was added dropwise to the above stirred solution followed by 3-hydrazinobenzoic acid (1 mmol), and the mixture was stirred overnight at RT. It was then diluted with 50 mL of cold water with vigorous stirring. The precipitate was collected by filtration, washed thoroughly with water several times, and then dried in vacuo overnight. The product was purified by flash column chromatography over silica gel eluting with 5 % methanol in dichloromethane. The pure product was obtained as a red to purple powder.
- N. Exemplary Compound 59-0125 where R¹ is methoxy, m is 1, the linker 30 is azo and Ar² is di(2-hydroxyethyl) amino, and related compounds having an azo

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linker can be prepared in a manner similar to that described by Alberti, G. et al. Chim Ind (Milan) (1974) 56:495-97.

O. Exemplary Compound 59-0124 and related, constrained analogs having the structure depicted below:

may be synthesized by reference to the methods described in Gorbulenko, N.V. et al. Dokl Akad Nauk Ukr SSR (1991) 5:117-23.

Related, constrained analogs having the structure depicted below:

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may be synthesized by reference to the methods described in the following publications: Chaurasia, M.R. & Sharma, A.J. Acta Cierc Indica Chem (1992) 18:419-22; Kandcel, Maymona M., in Phosphorus, Sulfur, Silicon, Relat Elem (1990) 48:149-55; Salem, M.A. & Soliman, E.A. Egypt J Chem (1985) 27:779-87, Garin, J. et al. Synthesis (1984) 6:520-22, or according to the representative procedure described in Ayyangar N. R. et al. Dyes and Pigments (1990) 13:301-10.

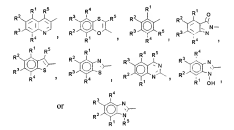
Claims

A method to treat a condition in a vertebrate animal characterized by a
deficiency in, or need for, bone growth or replacement and/or an undesirable level of
bone resorption, which method comprises administering to a vertebrate subject in need
of such treatment an effective amount of a compound of the formula:

wherein each of Ar^1 and Ar^2 is independently a substituted or unsubstituted phenyl, substituted or unsubstituted naphthyl, substituted or unsubstituted aromatic system containing a 6-membered heterocycle or a substituted or unsubstituted aromatic system containing a 5-membered heterocycle; and

L is a linker which spaces Ar¹ from Ar² at a distance of 1.5Å-15Å.

 $\label{eq:2.2} 2. \qquad \text{The method of claim 1 with the proviso that in the compound of} \\ 15 \qquad \text{formula (1), if } Ar^1 \text{ is}$



and L is

Ar2 cannot be

wherein

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5 R¹ is selected from the group consisting of:

H, OH, C1-C4 alkyl, C1-C4 alkoxy, C1-C4 alkylthio, halo and (C1-C12)alkyl-carbonyloxy;

R2 is selected from the group consisting of:

H, OH, halo, C1-C6 alkyl, C1-C6 alkenyl, C1-C6 alkoxy and (C1-C12)alkyl-carbonyloxy;

R3 is selected from the group consisting of:

H, OH, halo, C1-C6 alkyl, C1-C6 alkoxy, C1-C6 alkenyl and (C1-C12)alkyl-carbonyloxy;

R4 is selected from the group consisting of:

H, OH, halo, C1-C6 alkyl, C1-C6 alkoxy and (C1-C12)alkyl-carbonyloxy:

R5 is selected from the group consisting of:

H, halo, C1-C6 alkyl, C1-C6 alkoxy, -OC(=O)Me, phthalimide and (C1-C12)alkyl-carbonyloxy;

R6 is selected from the group consisting of:

20 H, OH, -NH₂, Cl-C4 alkyl and C1-C4 alkoxy;

R7 is selected from the group consisting of:

H, C1-C4 alkyl, (C1-C4)alkyl-carbonyl and (C7-C10)arylalkyl;

R8 is selected from the group consisting of:

H, OH, halo, -CF3, C1-C4 haloalkyl, C1-C4 alkyl, C1-C4 alkoxy,

5 -NHC(=O)Me and -N(C1-C4 alkyl):

R9 is selected from the group consisting of:

H, OH, halo, -CN, -NO₂, C1-C4 haloalkyl, C1-C8 alkyl, C1-C8 alkoxy, -NHC(=O)Me and -OC(=O)Me:

R¹⁰ is selected from the group consisting of:

H, OH, halo, -CN, -NO₂, C1-C4 haloalkyl, -CO₂H, C1-C12 alkyl, C1-C12 alkoxy, phenyl, C1-C12 alkenyl, (C1-C4)alkoxycarbonyl, -NHC(=O)Me, (C1-C4)alkylcarbonyl, (C1-C12)alkylcarbonyloxy and heteroaryl;

R11 is selected from the group consisting of:

H, OH, halo, C1-C4 haloalkyl, -CF3, C1-C4 alkyl, -NH2, C1-C4 alkoxy,

15 -NHC(=O)Me, C1-C4 alkenyl, (C1-C4)alkoxycarbonyl, (C1-C4)alkylcarbonyl, and (C1-C4)alkylcarbonyloxy;

R12 is selected from the group consisting of:

H, OH, $-NH_2$, C1-C4 alkyl, C1-C4 alkoxy and (C1-C4)alkylcarbonyl; and \mathbb{R}^{13} is selected from the group consisting of:

H, OH, halo, -NH2, C1-C4 alkyl, C1-C4 alkoxy -N(C1-C4)alkyl.

3. The method of claim 1 with the proviso that in the compound of formula (1), if Ar^{1} is

$$R^{a}_{m}$$
 Z Z X Ar^{1}

wherein R³ is a noninterfering substituent; m is an integer of 0-4;

each dotted line represents an optional π -bond;

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each Z is independently N, NR, O, S, CR or CR₂, where each R is independently H or alkyl (1-6C);

X is O, S, SO or SO2; and

L is a flexible linker.

5 then Ar² is not a substituted or unsubstituted 6-membered aromatic ring; if Ar¹ is

wherein Ra is a noninterfering substituent:

n is an integer of 0 and 5; and

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L is a flexible linker which does not contain nitrogen or is a constrained linker, then Ar² is not a substituted or unsubstituted phenyl or a substituted or unsubstituted naphthyl.

4. The method of claim 2 with the further proviso that in the compound of 15 formula (1), if Ar^{1} is

$$R^{a}_{m}$$
 Z Z Ar^{1}

wherein Ra is a noninterfering substituent:

m is an integer of 0-4;

each dotted line represents an optional π-bond.

each Z is independently N, NR, O, S, CR or CR₂, where each R is independently H or alkyl (1-6C);

X is O. S. SO or SO2; and

L is a flexible linker.

then Ar2 is not a substituted or unsubstituted 6-membered aromatic ring;

if Ar1 is

wherein Ra is a noninterfering substituent:

n is an integer of 0 and 5; and

L is a flexible linker which does not contain nitrogen or is a constrained linker, then Λr^2 is not a substituted or unsubstituted phenyl or a substituted or unsubstituted naphthyl.

5. The method of any of claims 1-4 wherein Ar1 is

$$R^{a}_{m} \longrightarrow Z \qquad (1a)$$
or
$$N \qquad (2a)$$

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wherein each Ra is a noninterfering substituent;

m is an integer of 0-4;

the dotted line represents an optional π bond:

Z is O, S, NR or CR₂ in formula (1) or is CR in formula (2) where each R is independently H or alkyl (1-6C), and

L is a flexible conjugating or nonconjugating linker or is a constrained linker.

 The method of claim 5 wherein L is a flexible conjugating or nonconjugating linker.

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The method of claim 6 wherein Z is NR

 The method of claim 7 wherein Ar² is a substituted or unsubstituted aromatic system containing a 5-membered heterocycle or is

wherein R^b is a noninterfering substituent and n is an integer of 0-5; and/or
L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or
-CONR- where R is H or alkyl (1-6C); and/or
the dotted line represents a n bond.

- The method of claim 7 wherein each R^b is independently halo, OR, SR,
 NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.
- m is 0; and/or

 each R^b is independently OR, SR or halo;

 where n=2 and at least one R^b is OR or SR; and/or

 L is -NHCO- or -CR=CR-.

The method of claim 7 wherein

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- The method of claim 7 wherein said compound is 59-0100, 59-103,
 59-104, 59-105 or 59-106.
 - The method of claim 6 wherein Z is S.
- The method of claim 12 wherein Ar² is a substituted or unsubstituted
 aromatic system containing a 6-membered heterocycle or is of the formula

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wherein R^b is a noninterfering substituent and n is an integer of 0-5, and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR- where R is H or alkyl (1-6C), and/or

5 the dotted line represents a π bond.

14. The method of claim 13 wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.

15. The method of claim 13 wherein
m is 0; and/or
each R* is independently OR, SR or halo;
where n=2 and at least one R* is OR or SR; and/or
L is -NHCO- or -CR=CR-.

- 16. The method of claim 12 wherein the compound is compound number 59-002, 59-0070, 59-0072, 59-0099, the benzothiazole counterpart of 59-0104, 59-0102, 59-0144, 59-0147, 59-0149, 59-0186, 59-0187, 59-0192, 59-0193, 59-0195, 59-0197, 59-0202, 59-0204, 59-0205, 59-0206, 59-0207, 59-0208, and 59-0210.
- The method of claim 16 wherein the compound is the benzothiazole counterpart of 59-0104, or is compound number 59-0147, 59-0205 or 59-0210.
- 25 The method of claim 6 wherein Z is CR or CR₂.
 - 19. The method of claim 18 wherein Ar² is

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wherein R^b is a noninterfering substituent and n is an integer of 0-5, and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR- where R is H or alkyl (1-6C); and/or

the dotted line represents a π bond.

- 20. The method of claim 19 wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.
 - The method of claim 6 wherein Z is O.
 - 22. The method of claim 21 wherein Ar2 is of the formula

15 wherein R^b is a noninterfering substituent and n is an integer of 0-5; and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR- where R is H or alkyl (1-6C); and/or

the dotted line represents a π bond.

- The method of claim 19 wherein each R^b is independently halo, OR,
 SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.
- The method of claim 21 wherein the compound of formula (1) is
 compound number 896-5005.

; and/or

- 25. The method of claim 5 wherein L is a constrained linker.
- The method of claim 25 wherein Z is S or NR; and/or wherein L is selected from the group consisting of

wherein Ar2 is

$$- \hspace{-1em} \bigwedge^{\mathsf{R^b}_m}$$

wherein Rb is a noninterfering substituent and m is 0-4.

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- 27. The method of claim 25 wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or comprises an aromatic system.
- 15 28. The method of claim 25 wherein the compound of formula (1) is 59-0124.
 - 29. The method of any of claims 1-4 wherein Ar¹ is of the formula

$$R^a$$
 N (3a)

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- wherein each R^a is independently a noninterfering substituent or is H; and Z is NR, S or O, wherein R is alkyl (1-6C) or H.

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30. The method of claim 29 wherein Z is S; and/or wherein Ar² is

wherein R^b is a noninterfering substituent and n is an integer of 0-5, and/or L is -N=N-, -N=CR-, -RC=CR-, -NRNR-, -CR₂NR-, -CR₂CR₂-, -NRCO- or -CONR- where R is H or alkyl (1-6C); and/or

31. The method of any of claims 1-4 wherein Ar1 is

$$R_m^a$$
 (4a)

wherein Ra is a noninterfering substituent;

m is an integer of 0-4;

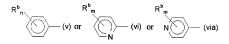
each dotted line represents an optional π-bond;

each Z is independently N, NR, CR or CR_2 , where each R is independently H or alkyl (1-6C) with the proviso that at least one Z is N or NR.

The method of claim 31 wherein Ar¹ is

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33. The method of claim 31 wherein Arz is



wherein each R^b is independently a noninterfering substituent, and n is 0-5 and m is 0-4; and/or

- 5 L is -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CO-, -NRNR-, -CR₂CR₂-, -NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-.
- The method of claim 33 wherein each R^b is independently halo, OR,
 SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.
 - 35. The method of claim 32 wherein each R^b is NR₂ or OR and m and n are 0, 1 or 2; and/or L is -CR=CR-,-N=N- or -NRCO-.
 - 36. The method of claim 35 wherein the compound of formula (1) is 59-0030, 59-0078, 59-0091, 59-0093, 59-0150, 50-0197, 59-0198, 59-0199 or 59-0480.
 - 37. The method of claim 31 wherein Ar_2 is substituted or unsubstituted quinolyl or naphthyl of the formula

20

wherein each Rb is a noninterfering substituent and m is 0-4.

- 38. The method of claim 37 wherein L is -N=N-, -RC=CR-, -RC=N-,
 -NRCO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CO-, -NRNR-, -CR₂CR₂-,
 -NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-; and/or
 wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃
 wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system and m is 0, 1 or 2.
- The method of claim 38 wherein the compound of formula (1) is 59-0089, 59-0090, 59-0092 or 59-0094.
 - 40. The method of claim 31 wherein Ar1 is

wherein each Ra is a noninterfering substituent and m is 0-4.

The method of claim 40 wherein L is -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CO-, -NRNR-, -CR₂CR₂-, -NRCR₂CR₂-, -NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-; and/or Ar² is

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wherein R^b is a noninterfering substituent and n is an integer of 0-5; and/or wherein each R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.

- 42. The method of claim 41 wherein the compound of formula (1) is 59-203, 59-285 or 59-286.
 - The method of claim 31 wherein L is a constrained linker.

44. The method of any of claims 1-4 wherein Ar1 is

$$\begin{array}{cccc}
R^{a}_{m} & z = z \\
z & & \\
z - z
\end{array}$$
(9a)

wherein each R^a is independently a noninterfering substituent; m is an integer of 0-4:

- each Z is independently N or CR, where R is H or alkyl (1-6C), with the proviso that at least one Z must be N and at least one Z must be CR.
- 45. The method of claim 44 wherein L is a flexible conjugating or nonconjugating linker; and/or

20 wherein Ar² is

$$R^b_n$$
 (v) or Z^b_{z-z} (vi)

wherein each Rb is independently a noninterfering substituent, and

in (vi) each Z is independently N or CR, where R is H or alkyl (1-6C), with the proviso that at least one Z must be a N and at least one Z must be CR.

46. The method of claim 45 wherein the compound of formula (1) is of the formula

$$\begin{matrix} R^a_{m} & & & R^b_{m} \\ & & & & \end{matrix}$$
 or
$$\begin{matrix} R^a_{m} & & & & \\ & & & & \end{matrix}$$

- 47. The method of claim 46 wherein L is -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CO-, -NRNR-, -CR₂CR₂-,
- 10 -NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-; and/or

wherein each R^a and R^b is independently halo, OR, SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system and each m and n is independently 0, 1 or 2.

- 15 48. The method of claim 47 wherein L is -NHCR $_2$ CR $_2$ NH-, m is 1 and R * is CF $_3$ para to L.
 - 49. The method of claim 48 wherein the compound of formula (1) is 59-0145, 59-0450, 59-0459 or 59-0483.

20

50. The method of any of claims 1-4 wherein Ar1 is

wherein each R^a is a noninterfering substituent; and n is an integer of 0 and 5, and wherein L is a flexible linker that contains at least one nitrogen; and/or

25

wherein Ar2 is of the formula

and L is -N=N-, -RC=CR-, -RC=N-, -NRCO-, -NRCR₂-, -NRCR₂CR₂-, -NRCR₂CCR₂-, -NRNRCR₂CCR₂-, -NRNRCR=CR-, -NRNRCOCR₂-, -NRNRCOCR=CR-, -NRNRCONR-, -NRNRCSCR₂-, -NRNRCSCR=CR-, -NRNRCONR-, -NRNRCSNR-, -NRNR-, -CR₂CR₂-, -NRCR₂CR₂NR-, -NRCR=CRNR- or -NRCOCR₂NR-.

- The method of claim 50 wherein each R^b is independently halo, OR,
 SR, NR₂, NO, NO₂, OCF₃ or CF₃ wherein R is H or alkyl (1-6C) or R^b comprises an aromatic system.
 - The method of claim 50 wherein L is -CR=CRCONRNR-, -CR=CRCSNRNR-, -CR2CONRNR- -CR2CSNRNR-, -NRNRCONR- or -NRNRCSNR- and/or

Rb is -NR2 and n=1 wherein Rb is in the para position.

- The method of claim 50 wherein R^a is -COOR and m is 1.
- 20 54. The method of claim 52 wherein the compound of formula (1) is 59-0045, 59-0095, 59-0096, 59-0097 or 59-0098.
 - 55. A pharmaceutical composition for use in a method to treat a condition in a vertebrate animal characterized by a deficiency in, or need for, bone growth replacement and/or an undesirable level of bone resorption which composition contains a pharmaceutically acceptable excipient and an effective amount of a compound of the formula set forth in any preceding claim.

- 56. A compound for use in preparing a composition for use in the treatment of a condition in a vertebrate animal characterized by a deficiency in, or need for, bone growth replacement and/or an undesirable level of bone resorption which method comprises administering said composition to a vertebrate subject, said compound set
- 5 forth in any preceding claim.

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Ar ¹ - lin 1.5 -	(I)	
Ar¹	Ar ²	
contains 5-membered heterocycle	substituted or unsubstituted benzene	II-A
contains 5-membered heterocycle	substituted or unsubstituted naphthalene	II-B
contains 5-membered heterocycle	contains 6-membered heterocycle	II-C
contains 5-membered heterocycle	contains 5-membered heterocycle	II-D
contains 6-membered heterocycle	substituted or unsubstituted benzene	II-E
contains 6-membered heterocycle	substituted or unsubstituted naphthalene	II-F
contains 6-membered heterocycle	contains 6-membered heterocycle	II-G
substituted or unsubstituted naphthalene	substituted or unsubstituted benzene	II-H
substituted or unsubstituted naphthalene	substituted or unsubstituted naphthalene	II-I
substituted or unsubstituted benzene	substituted or unsubstituted benzene	II-J

Figure 1

	CELLS		10/1/96					
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			READ 2		DUCTION AVE			
8-20	100.000		0 22	0.22.	0 18!	-0.99	-17 90	
	31 250	3.96	4 44	4 20	3.49	3.001	54.26	
	9.766	6 99	6.46	6 72	5.59	5.52	100.00	
	3.052	4 62	4 88		3.95	3.55	64 22	
	0 954	3.13	3 16		2.61	1.94	35 12	
	0 298	2.75	2 59	2.67	2.221	1 47	26.581	
	0.093	2.10	2.34	2.07	1.721	0.871	15.77	
	0.029	1 56	• 7:	: 63	1.36	0 43	7.80	
	0.0091	1 45	1 42	1 44	1.19	0.23	4.21!	
	0.0028	1 28	1.37	1.33	1.10	0.12	2.251	
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Figure 2

NNC#	MOL.WEIGHT	Concentration	% Response :
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50-0194	430.33		
50-0194		100.00 uM	-19.190
	1	31.25 uM	32.450
		9.77 uM	-14.240
		3.05 uM	-11.330
		953.67 nM	-12.790
	1	298.02 nM	-13.450
	+	93.13 nM	-12.290
	+	29.10 nM 9.09 nM	-9.440 -6.450
	+	2.84 nM	-8.130
	+	888.16 pM	-3.320
<u> </u>		333.10 (34)	
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1 - n - n - n - n - n - n - n - n - n -			1 1 1
50-0195	275.36		1 1
50-0195	215.30	100.00 uM	-4.630
1000193		31,25 uM	16.790
		9.77 uM	62.830
		3.05 uM	102.720
		953.67 nM	60.8601
		298.02 nM	32.450
	1	93 13 nM	19.340
		29.10 nM	17.220
	1	9.09 nM	5.640
		2.84 InM	4 840
		888.16 pM	5.640!
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50-0196	276.3	0	
50-0196	1 2,0,0	100.001uM	- 15.210
		31.25 uM	-8.560)
		9.77 uM	11.620
		3.05/uM	27.790
		953.67 nM	16.390
		298.02 nM	6.230
		93.13 nM	12.420
		29.10 nM 9.09 nM	12,630
		9.09 nM 2.84 nM	6.590
		866.18 pM	5.0601
		000.10 pM	2.0001

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50-0197		100.00	uM	-18.250	
		31.25		-14.980	
		9.77		4.040	
		3.05		93.790	
		953.87		205.530	
		298.02		242.920	
		93.13		195.890	
		29 10		115.320	
	-	9.09		85.630	
	+	2 84		54.380	
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9-0019	59-0019				
9-0019	59-0019				
		31.25			
		9,77			
				-17.470	
		3.05 ii		74 490	
				198.080	
		298.02	nM	258.340	
		298.02 93.13	nM nM	258.340 225.350	
		298.02 93.13 29.10	M M M	258.340 225.350 75.220	
		298.02 93.13 29.10 9.09	M M M	258.340 225.350 75.220 24.030	
		298.02 93.13 29.10 9.09 2.84	Mn Mc Mc	258.340 225.350 75.220 24.030 34.480	
		298.02 93.13 29.10 9.09	Mn Mc Mc	258.340 225.350 75.220 24.030	
		298.02 93.13 29.10 9.09 2.84	Mn Mc Mc	258.340 225.350 75.220 24.030 34.480	
		298.02 93.13 29.10 9.09 2.84	Mn Mc Mc	258.340 225.350 75.220 24.030 34.480	
		298.02 93.13 29.10 9.09 2.84	Mn Mc Mc	258.340 225.350 75.220 24.030 34.480	
		298.02 93.13 29.10 9.09 2.84	Mn Mc Mc	258.340 225.350 75.220 24.030 34.480	
		298.02 93.13 29.10 9.09 2.84	Mn Mc Mc	258.340 225.350 75.220 24.030 34.480	
CT ^N L C _O		298.02 93.13 29.10 9.09 2.84	Mn Me Mc	258.340 225.350 75.220 24.030 34.480	
N C C		298.02 93.13 29.10 9.09 2.84	Mn Me Mc	258.340 225.350 75.220 24.030 34.480	
N C C		298.02 93.13 29.10 9.09 2.84 888.18	MM TM TM TM TM TM	258.340 225.350 275.320 24.030 34.480 -3.740	
N C C	266.73	298.02 93.13 29.10 9.09 2.84 888.18 1	MM	258.340 223.350 75.220 24.030 34.480 -3.740	
\$-0070 \$-0070	266.73	298.02 93.13 29.10 9.09 2.84 888.18	MM IM IM IM IM IM IM IM IM IM	258.340 225.350 75.220 24.030 34.480 -3.740	
\$-0070 \$-0070	266.73	298.02 93.13 29.10 9.99 2.84 888.18 9.00 100.00 31.25	nomina no	258.340 225.350 75.220 24.030 34.480 -3.740 -16.510 -16.640 -0.270	
\$-0000 \$-0000	266.73	298.02 93.13 29.10 10.00	1944	258.340 225.350 75.220 24.030 34.480 -3.740 -18.510 -18.040 -0.270 95.440	
→ 0020 → 0020	266.73	298.02 93.131 29.100 9.090 2.84 888.1819 100.0010 31.25 9.7710	IN I	258.340 225.350 75.220 24.030 34.480 -3.740 -16.510 -16.640 -0.270	

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. 29.10 nM	37.870
9.09 nM	24.820
2.84 nM	20.500
888.18 pM	13.310

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-0021	284.72	100.00 uM	-16.310
9-0021	+	31.25 uM	+12.850
			84.130
		3.05 uM	89.940
		953.87 inM	1 65.750
	1	298.02 nM	33.940
		93.13 nM	22.560
		29.10 nM 9.09 nM	13.910
	-ii-	2.84 nM	33.270
		888.18 pM	15.5001
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9-0022	268.37		
59-0022		100.00 uM	7.250
		31.25 uM	-5.010
	1	9.77 uM	-0.270
	1	3.051uM 953.571nM	3.060
		298.021nM	-1.800
		93.13inM	-0.200
	- -	29.10 nM	-3.270
		9.09inM	1 130
		2.84 inM	2.590
		888.18 DM	2.460
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	239.28		
59-0023	239.20	100.001uM	-12.720
59-0023		31.251uM	33.140
		9.77 uM	56.5001
	i	3.051uM	29.550
		953.67 nM	25.3601
	-	298.02 inM	15.700
		93.13 nM	7.380
		29.10InM	-9.710
		9.09 InM	1 000
	1		4 520
		888.18 pM	-0.010

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59-0024			
35-0024	220.28		1 1
'		1.	
59-0025	224.31		
59-0025		100.00 luM	-25.590
		31.25 uM	14.150
		9.771uM	50.890
		3.05 uM	57,880
	1	953.67 InM	38.900
		298.02 nM	28.530
		93.13 nM	19.660
		29.10 nM	17.490
		9.09 nM	-0.600
		2.84 nM	4 190
		888.18 pM	4.870
59-0026	248.29		
59-0026	248.29	100.00 uM	
		31.25 uM	-29.830
		9.77 uM	
		3.05 uM	-10.470
		953.57 nM	46.220
		298.02 nM	107.780
		93.13 nM	36.850
		29.10inM	28.720
		9.09 nM	8.520
		2.84 nM	-1.240
		888.18 pM	4.020

NH IFH			
, hard			
59-0027	250.30		
59-0027	250.30	100.00 uM	
		31.25 uM	89.810
		9.77 uM	54.870
		3.05 uM	44 940 23 780
		953.67 inM	8.380
		298.02 nM	6.330
		93.13 nM	7.360
		29.10 nM	3.380
		9.09inM	-1.620
		2.84 nM	-3.570
		888,18 pM	-0.720
N N N N N N N N N N N N N N N N N N N			
59-0028 59-0028	226.28	100.00 uM	-28.750
		31.25 uM	-28.750I -16.740I
		9.77 JuM	29.550
		3.05 JuM	100.580
		953.67 nM	54 940
		298.02 nM	31.340
		93.13 nM	7 500
		29.10 nM	7.500
	,	9.09InM	7 880
		2.84 nM	3 140
		888.18 pM	4 670:

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59-0029		i	i
59-0029	249.27		
		100.00 uM	-15.160
		31.25 uM	41.940
		9.77 uM	36.630
		3.05 uM	7.120
		953.67 nM	21.880
		298.02 nM	15.540
		93.13 nM	1.810
		29.10 nM	1.370
		9 09 nM	12.140
		2.84 mM	4 230
		888.18 pM	9.040
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59-0030 A		1	1 1
59-0030 A	233.28		
35-0030 F(100.00 uM	-27.970
		31.25 uM	-22.830
		9.77 uM	-5.420
		3.05 uM	57.280
		953.67 nM	72.620
		298.02 InM	53.000
		93.13 InM	29.990
		29.10 nM	14 630
		9.09 nM	3.870
		2.84 nM	6.970
		888.18 pM	1.810
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	231 30	31.25 uM 9.77 uM 3.05 uM	-17 810 - 20.840 87.380
59-0031	231 30	31.25 uM 9.77 uM 3.05 uM 953.67 nM	-17 810 - 20.840 87.380 49.320
59-0031	231 30	31.25 uM 9.77 uM 3.05 uM 953.67 nM 298.02 nM	-17 810 - 20.840 - 87.380 - 49.320 - 43.110
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59-0032	248.29		
59-0032		100.00 uM	-7.780
		31.25 luM	40.750
		9 77 JuM	42.820
		3.05 uM	25.700
		953.67 nM	31.170
		298.02 nM 93.13 nM	34.410
		29.10 nM	3.570
		9.09 nM	4.320 -10.000
		2.84 nM	5.650
		888.18 pM	11.990
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59-0033	248.29		1 1
59-0033		100.00 uM	-28.180
		31.25 uM	-11.590
		9.77 uM	55,3001
		3.05 uM	49.710
		953.67 nM	47.410
		298.02 nM	0.250
		93.13 nM	7.9801
		29.10 nM	-8.940
		9 09 nM 2.84 nM	-7.6301
		888.18 pM	-0.4001 -5.9801
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59-0034		1	
0.0004	268.34		
9-0034		100.001uM	-28.51
		31.25 uM 9.77 uM	24
		3.05 uM	73 58
		953.67 nM	20.09
		298.02 nM	16.87
		93.13 nM	15.23
		29.10 nM	28.83
		9 09 inM	9.08
		2.84 InM 888,18 IpM	_23.02 -0.32

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59-0035	291.36	1	
59-0035		100.00 luM	-14.92
		31.25 uM	29.17
		9.77 uM	15.87
		3 05 luM 953 67 lnM	18.81
	<del></del>	298.02 nM	3.88
		93.13 InM	8.15 3.22
		29 10 nM	-10.03
		9.09 nM	15.58
		2.84 inM	-3.56
		888.18 pM	-7 13
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9-0036	262.31		
9-0036		100.00IuM	-0.98
		31.25 uM	-3.25
		9.77 uM	4.54
		3.05 uM	-1.95
		953.67 inM	0.32
		298.02 nM 93.13 nM	-6.491
		29.10 nM	-17 191 -0.561
		9.09 nM	-5.52
		2.84 nM	-9.4
		888.18 pM	-16.53
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		31.25luM	-11 99
		9.77 luM	-10.03
		3.05 uM	-19 11:
		953.67 InM	-9.4
	- 1	298.02 InM	2.27
		93.13 inM	-2.9
		29.10InM	-10.69
		9.091nM 2.841nM	2.59
		888.18 pM	0.66

59-0038			
59-0038	291.36		
		100.001uM 31.25 uM	-23.4301
	<del></del>	9.77 uM	-8.3901 -0.1001
		3.05 uM	-2.880
		953 67 InM	-2.240
		298.02 nM	3.900
	<del></del>	93.13 nM	8.350
		29.10 nM 9.09 nM	1.150)
		2.84 nM	8.960
		888.18 pM	-0.3801
OH NNNN SNNSN			
59-0039	312.35		
9-0039		100.00 luM	14.170
		31.25 uM	7.520
		9.77 uM	1.940
		3.05 uM 953.67 inM	-3.140
		298.02 InM	-7.770 -5.980)
		93.13InM	-8.820
		29.10InM	-2.390
		9.09 inM	-16.5801
		2.84 InM	-4 480
		888.18 pM	-0 450:
9-0040	290.37		
9-0040	230.371	100.00 uM	-20 400
		31.25iuM	-17 310
		9.77 uM	-8 110
		3 05 luM	32.180
		953.67 nM	36.180
		298.02 nM	17.440
		93.13 InM 29.10 InM	2.040
		9.091nM	10.350
		2.84 nM	6 960

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59-0041			
59-0041	501 90		
		100.00 uM 31.25 uM	-18 37:
		9.77 UM	-17.33: -5.111
		3.05 luM	3.31
		953.87 nM	-0.77
		298.02 nM	-1.58
		93.13 nM 29.10 nM	3.55
		9.09 nM	-11.24
		2 84 nM	0.25
		888.18 pM	2.02
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59-0042	281.36		i
59-0042		100.00 uM	183.51
		31.25 UM	-7.67
		9.77 uM	9.41
		3.05 uM 953.67 nM	0.75
		298.021nM	6.11I 3.82I
		93.13 nM	2.54
	1	29.10 InM	4.07
		9.09 nM	-9.73
		2 84 inM	-0.02
		888.181pM	18.37
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9-0043	280.29		1 1
9-0043		100.00 uM	20.66
		31.25 uM	7.4
		9 771 UM 3.05 UM	-1.29
		3.05 uM 953.67 nM	-2.31!
		298.02 nM	1.54
	-	93.13 nM	1.52
		29.10 nM	2.79
		9 09 nM	0.27
		2.84 nM 888.18:pM	8 92

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59-0044	341.21		i !	
59-0044		100.00 uM	7 381	
		31.251uM	7.38	
		9.77 iuM		
		3.05 uM	12.49	
		953.67 InM	0.52	
		298 02 nM	6.11	
		93.13InM	-1.54	
		29.10inM	19.14	
		9.09InM	7.13	
		2.84 nM	-2.06	
	i-	888.18 pM	-2.06 5.84	
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59-0045	283.33	!	i :	
59-0045	1.	100.00 luM	52.37	64 460
		31.25 iuM	148.43	192.960
		9.771uM	204.47	422.540
	-	3.05 luM	280.3	437 020
		953 87 InM	254 82	410.890
		953 87 InM 298.02 InM	254 82 218 21	
		298.02 inM	218.21	266 090
		298.02 inM 93.13 inM	218.21 196.98	266 090 183.730
		298.02 inM 93.13 inM 29.10 inM	218.21 196.98 96.06	266 090 183.730 80.440
		298.02 inM 93.13 inM 29.10 inM 9.09 inM	218.21 196.98 96.06 67.35	266 090 183.730 80.440 55.530
		298.02 inM 93.13 inM 29.10 inM	218.21 196.98 96.06	266 090 183.730 80.440 55.530
		298.02 inM 93.13 inM 29.10 inM 9.09 inM	218.21 196.98 96.06 67.35	266 090 183.730 80.440 55.530
		298.02 inM 93.13 inM 29.10 inM 9.09 inM	218.21 196.98 96.06 67.35	266 091 183.73 80.44 55.53
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9-0046		298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.64 inM	218.21 196.98 96.06 67.35	266 090 183.730 80.440 55.530
59-0046	389 37	298.02 inM 93.13 inM 29 10 inM 9 09 inM 2 84 inM	218.21 196.98 96.06 67.35 52.99	266 090 183.730 80.440 55.530
59-0046	389 37	298.02:nM 93.13:nM 29.10:nM 9.09:nM 2.64:nM	218.21 196.98 96.06 67.35 52.99	266 090 183.730 80.440 55.530
59-0046	389 37	298.02:nM 93.13:nM 23.10:nM 9.09:nM 9.09:nM 2.84:nM	218.21 196.98 96.06 67.35 52.99	266 090 183.730 80.440 55.530
59-0046	389 37	298.02:nM 93.13:nM 29.10:nM 9.09:nM 2.84:nM	218.21 196.98 96.06 67.35 52.99 79.33 2.24, -1.87 -6.18	266 090 183.730 80.440 55.530
59-0046	389 37	298 02 rM 93.13 rM 99.13 rM 99.01 rM 9.00 rM 2.04 rM 100.00 ruM 31.25 ruM 3.05 ruM 3	218.21 196.98 96.06 67.35 52.99	266 090 183.730 80.440 55.530
59-0046	389 37	298 02 rbM 93.13 rb 93.13 rb 29 10 rb 9 09 rb 100 00 rb 100 00 rb 100 00 rb 110 00 rb 110 00 rb 110 00 rb 110 00 rb 110 r	218.21 196.98 96.06 67.35 52.99 79.33 2.24, -1.87 -6.18	266 090 183.730 80.440 55.530
59-0046	389 37	298 02 rbM 93.131nM 29.101nM 9.091nM 2.041nM 2.041nM 100.00 uM 31.231uM 3.051uM 30.501uM 39.53.371nM 298.021nM 39.311nM	218.21 196.98 96.06 67.35 52.99 79.33 2.24 -1.87 -6.18	266 090 183.730 80.440 55.530
59-0046	389 37	298 02 inM 93.13 inM 29.10 inM 9.09 inM 2.04 inM 100.00 iuM 31.25 iuM 9.77 iuM 3.05 iuM 953.37 inM 953.37 inM	218.21 196.98 90.06 67.35 52.99 79.33 2.24, 1.187 6.18 0.001	266 090 183.730 80.440 55.530
59-0046	389 37	298 02 rnM 93.131nM 29.101nM 9.001nM 9.001nM 100.00 uM 11.201nM 100.00 uM 31.25 uM 31.25 uM 30.5 uM 953.87 rnM 298.02 rnM 298.02 rnM 9.001nM 298.02 rnM 298.02 rnM 29	218.21 196.98 96.06 67.35 52.99 79.33 2.24 -1.87 -6.18 -0.001 -3.63 -0.84	410.89C 266.09C 183.733 80.44C 55.533 44.16C
59-0046	389 37	298 02 inM 93.13 inM 29.10 inM 9.09 inM 2.04 inM 100.00 iuM 31.25 iuM 9.77 iuM 3.05 iuM 953.37 inM 953.37 inM	218.21 196.98 90.06 57.35 52.99 79.33 2.24 1.87 4.18 0.001 1.3.63	266 090 183.730 80.440 55.530

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59-0047	303.37	
59-0047	100.00luM	-6.73
	31.25luM	10.38
	9.77 uM	-6.16
	3.05/uM	-1.39
	953.67 nM	-10.11)
	298.02 nM	4 49
	93.13 nM	-7.28
	29.10 nM	-12.341
	9.09 nM	-3 08
	2.84 nM	-2.26
	888 18 pM	-5.34
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59-0048	384.50	
59-0048	100.001uM	-6.73
	31.25 uM	0.27
	9.77 uM	-5 61
	3.05 uM	-2.26
	953.67 nM	-12.89
	298.02 nM	-1.69
	93.13 nM	4.77
	29 10 lnM	-8.14
	9.09inM	-3.92
	2.84 InM	-11.2
	888.18 pM	-4.77
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59-0049	251.29	1
59-0049	100 00 luM	4 49
	31.25 uM	. 0
	9 77 luM	4.77
	3.05 uM	1.96
	953.67 inM	8.69
	298.02:nM	-5 04
	93.13 nM	-2.24
	29.10inM	1.69;
	9 09 nM	-4 49
	2.84 inM	2.24
	888.18 pM	-0.31

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59-0050	303.36	1 1
59-0050	100.00 uM	45.79
	31,25 uM	10.02
	9 77 iuM	11.29
	3.05 uM	4 68
	953.67InM	6.92
	298.021nM	-5.65
	93.13 nM	1.69
	29.10 nM	-7.57
	9.09 nM	-12.05
	2.84 nM	-13.63
	888.18 pM	5.2
59-0051	251.35	
59-0051	100.00 uM	32.36:
	31.25 uM	-18.42
	9.77 uM	-0.55
	3.05 uM	-13.94
	953.67 nM	12.02
	298.02 nM	-14 59
	93.13 nM	-7 55
	29.10 nM	-114
	9.09 nM	-14 91
	9.09 nM 2.84 nM	-14 91 -10 74

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59-0052			i
59-0052	393.28		
39-0052		0.00 uM	-21 52:
		1.25 uM	-13.32:
		9.77 uM !	-21.311
		3.05luM	-11 08
	95	3.67 InM	-20.66.
	1 29	3.02 InM	-17.14/
	9:	3.13 nM .	-16.491
		9.10 nM	-11.4
		9.09 nM	-10.74
		2.84 inM	-11,081
		3.18 pM	-14 59
59-0053	354 41	i	
59-0053		100:uM	-17 14
	31	251uM	-21 31'
		771uM	-9 47
		.05:uM	-11 08:
		67(nM	-0.83
		.02 nM	-114
		.13/nM	-9 47
		.10inM	-19.72
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		.84:nM	-10.09
		.181pM	
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59-0054		1	1 1
59-0054	236.28		
-		100.00 LM	-20 04
		31.25 UM	-6.95
		9 77 luM	83
		3.05 luM 953.57 lnM	-3.37
-		298.02 inM	-2.4
		93.13 nM	-0.00
		29.10 nM	-0.99
1		9.09 nM	
		2.84 inM	5.92
		888.18 pM	-2.17
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59-0055	425.51		1
59-0055		100.001uM	-13.76
		31.25luM	-9.51
		9.77 uM	-2.02
		3.05 JuM	3.24
		953.67 InM	-6.27
		298.02 InM	4.05
		93.13 InM	-1.62
		29.10InM	-7 49
		9.09 inM	-7 09
		2.84:nM	
		2.84 inM	-3.04
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59-0056	512.34	- 1	1 1
9-0056		100.001uM	-1 42
		31.25 uM	4 871
	-	9.77 uM	0.18
		3.05 UM	3.84:
		953.67InM	-5.07
		298.02 nM	-7.29
		93.13 nM	
		29.10inM	0.0011
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		9.091nM 2.84 nM	-1 02 -3.85

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59-0057	į į	
59-0057	100.00 uM	-24.150
	31.25 uM	-24.300)
	9.77 uM	-5.980
	3.051uM	-11.500
	953.67 nM	-13.000
	298.02 nM	-6.280
	93.13 nM	-12.550
	29.10 nM	-8.870
	9.09 lnM	-8.5201
	2 84 nM	-16.290
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59-0058		
59-0058	100.00 uM	4 170
	31,25 uM	7.620
	9.771uM	-1.790
	3.051uM	-7.320
	953.67 InM	-1.940
	298.02:nM	-6.870
	93.13 nM	-1 490
	29.10InM	-8 370
	9.09 nM	-5 080
	2.84 nM	-12 400
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9-0059	i	
9-0059	100 00 iuM	-18 770.
	31.25!uM	-16 140
	9.77:uM	-3 090!
	3.05 uM	0 150
	953.67 nM	6.010
	298.02 nM	-1.910
	93 13 nM	-1.760
	29.10 nM	9.100
	9.09 nM	-8.220
	2 84 nM	-5 7201
	2.041104	-5 /20

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9-0060		
9-0060	! 100 00iuM	4 250
	31.25 uM	-14.520
	9 77 luM	1.030
	3.05 uM	-1.180
<u> </u>	953.67 nM	-13,200
- <u> </u>	298.02 nM	-0.740
	93.13)NM	-3.670
-	29.10 nM 9.09 nM	-7.340
	2.84 nM	-1.310 0.290
	4.041mM	0.290
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9-0061		
9-0061	100.00 uM	1 -17.890
	31.25 uM	-18.770
1	9.77 uM	-17.170
	3.05 uM	-14.080
	953.67 InM	-17.020
	298.02 nM	-7 190
	93.13 nM	-1.910
	29.10 nM	-0 4401
	9.09 nM	-6.010 -4.560
	2.84 nM	4 560
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9-0062		1
9-0062	100.001uM	-13.940
	31.25 uM	-12.910
	9.77 uM	-4.560
	3.05 uM	-4.540
	953.67 inM	-5.900
	298.02 nM 93.13 nM	-1 620

	9.09inM	8.070
	2.84 inM	0 440.
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59-0063	1 !	8 1
59-0063		<u> </u>
	100.00 uM	-2.510
	31.25 uM	-6.1301
	9.77 UM	-8.950
	3.051UM	-8.020
	953.67 inM	-6.010
	298.02 nM	-2.520
	93.13InM	-5.810
	29.10 nM 9.09 nM	-3.450
	2.84 nM	-4 390
	2.84 nm	-6.280
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59-0064		
59-0064	100.00 UM	-23.090
	31.25luM	-21.040
	9.771uM	78 400
	3.05 luM	155.220
	953.67 InM	113 120:
	298.02 nM	30.6401
	93.13 nM	15.240
	29.10 inM	22.150:
	9.091nM	-0.7701
	2.84 InM	4.410
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9-0065		·
3-0003	100.00 uM	-2.0301
	31.25 uM	-2 980
	9.77 uM	-15.240:
	3.05 uM	-15.4001
	953.67 InM	-15.240
	298.02 nM	-10.520
	93 13 InM	-13.830
	29.10 nM	-5.810
	29.10 nM i 9.09 nM 2.84 nM	-5.810

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59-0066	1 1	1 1
59-0066	100.00 juM	
1	31.25 uM	10.060
	9 77 UM	2.580
	3.05 uM	10.850
	953.67 InM	14.610
	298.02 nM	0.950
	93.13 nM	3.7601
	29.10 mM	1.730
	9.09 nM	-2.820
	9.09 nM 2.84 nM	-2.820
	2.04 INM	-3 920
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59-0067		1
59-0067		
-	100.00 uM	-24.040
	31.25 uM	-24.690
	9.77 uM	-1.450
	3.05 uM	60.900
	953.67 nM	133,860
	298.02 nM	75.330
	93.13 nM	26,760
	29.10 nM	20.070
	9.09 nM	4 980
	2.84 nM	4 450
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9-0068		!
9-0068	100.00 luM	-22,130
	31.25 uM	-7 880
	9.77 uM	93.900
	3.05 uM	81.060
	953.67 InM	22.330
	298.02 InM	17.300
	93.131nM	8.460
	29.10!nM	-3.530
	9.09 nM	4 230
	2.84 nM	─6 140

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59-0069 59-0069		
19-0069	100.00 uM	5.490
	31.25 uM	9.670
	9.77 uM	16.090
	3.05 uM	-7.180
	953.67 nM	-2.840
	298.02 nM	-3.710
	93.13 nM	-11,180
	29.10 nM	-5.790
	9.09 nM	-7.180
	2.84 nM	4.750
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59-0070		1
59-0070	100.00 iuM	-25.930
	31.25 uM	-23.000
	9.77 UM	35.060
	3.05 iuM	214.2801
	953.67 nM	158 530
	298.02 nM	72.890
	93.13 nM	20.9401
	29.10InM	7.760!
	9.09 nM	7.5901
	2.84 inM	-8 400:
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59-0071		
59-0071	100.00 uM	-18 650
	31.25 uM	-15.540
	9.77 uM	17 060
	3.05 uM	178 090
	953.67InM	78 070
	298.02 nM	31.260
1	93.13 nM	18.410
	29 10 nM	4.870
	9.09 nM	
	2.84 nM	
	2.041nM	

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59-0072		
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50.0012	100 00 uM	-19 750
	31.25 uM	-18.650
	9.77 uM 3.05 uM	-18 430
	953.67 nM	-15.770
	298.02 nM	9.970
	93.13 nM	74.740
	29.10 nM	175.430
		213.580
	9.09 nM 2.84 nM	164.320
	888.18 pM	119.100
	008.18 IPM	60.770
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59-0073 59-0073		
58-00/3	100 00 iuM	-3.010
	31.25 uM 9.77 uM	4.830
	3.05 uM	-9.660 -4.680
	953.67 InM	-6 500
	298.02 nM	-2.510
	93.13InM	7 140
	29.10 nM	0.97
	9.091nM	-5.5
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59-0074	100 00 iuM	-2.85
	31.251uM	2 14
	0.771.44	
	9 77 uM	4.85
	3.05 uM	-3.5
	3.05 uM 953.67 nM	-3.5 -4.85
	3.05 uM	-3.5
	3.05 uM 953.67 nM 298.02 nM	-3.5 -4.85 9.95
	3.05 iuM 953.67 inM 298.02 inM 93.13 inM	-3.5 -4.85 9.95 -4.47

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59-0075		
59-0075	100.001uM	-0.68
	31.25 uM	-10 16
	9.77 UM	-5.35
	3.05 uM	-8.5
	953.67 nM	-0.85
	298.02 nM	5.97
	93 13 nM	0.97
	29.10 inM 9.09 inM	-2 35
	2.841nM	0.32
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59-0076		
59-0076	100.00luM	-19.121
	31.25 uM	9.29
	9.77(uM	10.83
	3.05luM 953.57lnM	22.43
	298.02 nM	3.47
	93.13InM	19.93
	29.10InM	10.63
	9.09inM	14.28
	2.84 nM	11 3
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59-0077		
9-0077	100.00 uM	-20.96
	31.25 uM	-16.23
	9.77luM	-10.58
	3.05 uM	-11.96
	953.67 InM	-19.44
	298.02 nM	-17.3
	93.13 InM	-13.79
	29.10 nM 9 09 nM	-15.62
	9 09/nM 2 84/nM	-14 09
	2.04-NM	144

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59-0078		
	100.00 JuM	-25.540
	31.25 uM	-22.560
	9.77 uM	71.5301
	3.05 uM	207.960
	953 67 nM 298.02 nM	379.230
	93.13 nM	241.460
	29 10 mM	84.020
	9.09 nM	50.350
	2.84 inM	56.600
	0.80 nM	92.520
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59-0079		
59-0079		
370073	100.00 uM	-34.980
	31.25 uM 9.77 uM	-21.390
	3.05 uM	1 37.200
	953.67 nM	69.010
	298.02 nM	64.000
	93.13 nM	46 490
	29.10 nM	30.310
	9 09 nM	33 490
	2.84 nM	29.760
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9-0080		
9-0080		1 1
	100.001uM 31.251uM	5 3901
	9.77 uM	5.560
	3.05 uM	2 440
	953 67 InM	-5.030
	298.021nM	7 660
	93.13 InM	-3.630
	29.10InM	3.650
	9 09 nM 2.84 nM	1.050
	2.64 INM	6.940
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9-009v	' 1	1

59-0081		_
1	300.00 uM	62 840
	9.77 uM	11 300 -8 6701
	3.05 uM	2 440
	953.67 nM	-5.200
	298.02 nM	-2.080
	93.13 nM	1.220
	29.10 nM	-2.250
	9.09 nM	1.050
	2.84 nM	-3.300
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59-0082		
59-0082	100.00 uM	111.79
	31.25 uM	62.88
	9.77 iuM	32.36
	3.05 uM	9.11
	953.67InM	-10.62
	i 298.02 inM	-1.86
	93.13 nM	-6.89
	29.10 nM	-3.91
	9.09 nM	2.22
	2.84 nM	16.36
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59-0083		1 1
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59-0083	100.00 iuM	48.93
	31.25 uM 9 77 uM	40.91
	3.05 UM	25.85
	953.67 InM	8.55
	953 67 InM 298 02 InM	8.55
	953-67 InM 298-02 InM 93-13 InM	8.55 3.9 2.05
	953 67 InM 298 02 InM 93.13 InM 29.10 InM	8.55 3.9 2.05 7.99
	953 67 inM 298 02 inM 93.13 inM 29.10 inM 9 09 inM	8.55 3.9 2.05 7.99
	953 67 InM 298 02 InM 93.13 InM 29.10 InM	8.55 3.9 2.05 7.99
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	953 67 inM 298 02 inM 93.13 inM 29.10 inM 9 09 inM	8.55 3.9 2.05 7.99
C C OH	953 67 inM 298 02 inM 93.13 inM 29.10 inM 9 09 inM	8.55 3.9 2.05 7.99
59-0014	993.07/nb/ 299.02/nb/ 99.313/nb/ 29.10/nb/ 29.10/nb/ 29.10/nb/ 29.10/nb/ 29.10/nb/	3.91 3.91 2.051 7.799 3.391 3.391
C C OH	99.3 67 / m/s 298 02 / m/s 99.3 13 / m/s 29.1 01 / m/s 9 0.9 / m/s 2 2 8 4 / m/s	3.9 3.9 2.05 7.799 3.35 3.35 3.35
59-0014	993 67 (nb) 299 02 (nb) 99 3 13 (nb) 29 9 10 (nb) 29 9 10 (nb) 29 9 10 (nb) 2 9 10 (nb) 2 9 4 (nb) 2 8 4 (nb) 100 00 (ub) 31 25 (ub)	3.9 2.05 7.99 3.91 3.91 3.91 3.35 3.35
59-0014	99.3 67 / m/s 298 02 / m/s 99.3 13 / m/s 29.1 01 / m/s 9 0.9 / m/s 2 2 8 4 / m/s	3.9 3.9 2.05 7.799 3.35 3.35 3.35

	953.67 InM	21.700
	298.02 nM	5 9001
	93.13 inM	4 870
	29 10 InM	1 -10 920
	9.09 nM	10.080
	2 84 nM	-2.080
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59-0085 59-0085		
59-0085	100.001uM	17.070
	31.25 LuM	41.890
	9.77 LuM 3.05 LuM	18.500
	953.67InM	20.340
	298.02 nM	22.4901 8.090
	93.13 nM	11 790
	29.10inM	1,240
	9.09 nM	-0.7601
	2.84InM	5.940
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59-0086	1 1	1
9-0086	100.00 uM	30.750
	31.25 uM	31.190
	9.77 uM	14.790
	3.05 iuM	13,500
	953.87 nM	14.080
	298.02 nM	3.940!
	93.13InM	9.370
	29.101nM	-2.510
	9.091nM	-5.040
	2.841nM	1.530
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9-0087		
9-0087	100.00 uM	10.660
	31.25 uM	: 11.080
	9.77 UM	3.100
	3.05/uM	-1.320
	953.67 InM	17.070
	298 02 InM	7.950
	93.13!nM	-4.460
	29 10 inM 9.09 inM	4.510
	2 84 nM	-0.470
		9.660

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59-0088	1 1	į l
59-0088		
35-0088	100.001uM	
	31.25 uM 9.77 uM	
	3.05 uM	
	953.67 nM	
	298.02 nM	
	93.13 nM	
	29.10 nM	
	9.09 nM	
	2.84 nM	
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59-0089		
59-0089	100.00 luM	60.09
	31.25 uM	118.25
	9.77 uM	65.84
	3.05 uM 953.67 nM	36.11
	298.02 nM	37.96 18.42
	93.13 nM	6.33
	29.10 InM	13.58
	9.09 nM	0.75
	2.84 nM	-5.77
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9-0090		
9-0090	100.00 uM	32.77
	31.25/uM	24.63
	9.77iuM	19.5
	3.05iuM	41.31
	953.67 InM	9.81
	: 298.021nM	-1 76
	93.13InM	3 53
	29.10InM	2.95
	9.09 nM	2.95
	2.841nM	7.8
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9-0091	1 1	
9-0091	100.00 uM 31.25 uM	0.261

	9 77 uM	95 94
	1 3.05 uM	87 71
	953.67 inM	44 17
	298.02 nM	30.20
	93.13 nM 29.10 nM	23.071
	9 09 nM	21.65
	2.84 inM	10.95
	2.04/////	20.92
59-0092		
59-0092	100.00 uM	
374031	100.00/uM 31.25/uM	-11.56
	9.77 uM	17 84 50.19
	3.05 uM	25.84
	953.87 inM	14.4
	298.02 nM	6.77
	93.13 nM	8.62
	29.10InM	2.22
	9.09 mM	6.38
	2.84 nM	11
59-0093		
59-0093	100.00 uM	-11.67
	1 31.25 uM	15.02
	9.77 uM	35.44
	3.05 uM	29.89
	953.87 nM	22.881
	298.02 nM 93.13 nM	19 561 5 18
	29.10 nM	7.39
	9.09 nM	4.56
	2.84 nM	5.9
59-00p4		
59-0094	100.00	17.00
33-03-	100.00 uM 31,25 uM	-17 69: - 45.15
	9.77 uM	24.97
	3.05 uM	19.81
	953.67 nM	9.35
	298.02 nM	1.36
	93 13 nM	9.24
	29.101nM	-0 48
	9.09 nM	6.18
	2.84 nM	1 611

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59-0095 59-0095		
159-0095	100.001uM	44
	31.25 uM	47.6
	9.77 uM	12 7
	3.05 uM 953.67 nM	21 4
	298.02 InM	15.0
	93.13 nM	10.22
	29 10 nM	20.3
	9 09 nM	10 9
	2 84 nM	9.2
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59-0096		1
59-0096	100.00 uM	413.05
	31.25 uM	287.23
	i. 9.77 uM	137.36
	3.05 uM	78.5
	953.67InM	49.13
	298.02 nM	50.68
	93.13 nM	47.95
	29.10 nM	26.28
	9.09InM I	18.75
	2.84 nM	22.17
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9-0097	!	
	100.00 uM	77.47
	31.25 uM	201.9
	9.77 uM	160.93
	3.05 uM	81 44
	953.67 nM	47.78
	298.02 inM	51.54
	93.13InM 29.10InM	34 64
	40.101.00	43.18
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59-0098	1 1	
59-0098	100.00 uM	
	31.25 uM	-1.3E
	9.77 uM	156.89
	3.05 uM	164.69
	953.67 nM	96.94
	298.02 InM	68.2
	93.13 InM	5
	29 10 InM	51.8
	9.09 nM 2.84 nM	41 2
	2.64 INM	33.43
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59-0099		1 1
59-0099	100.00 uM	13.040
	31.25 iuM	13.040i 56.880:
	9.77 uM	119.340
	3.05 uM	237.420:
	953.67 nM	285 4401
	298.02 nM	164.610
	93.13 nM	123.300
	29.10 nM	69.2401
	9.09 nM	44.500
	2.84 nM	47.390
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9-0100	1 1	
9-0100	100.00 uM	10,000
	31.25 uM	-10 0201 -10.7301
	9.77 uM	30 340
	3.05 uM	114.410
	953.67 inM	77 540
	298.02 nM	40.290
	93.13InM	35.7301
	29 10 nM	28.290
	9.09 nM	17.480
	2.84 nM	11.470
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9-0101		
19-0101	100.001uM	26 370

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		31.25.uM	12 440
		9 77 iuM	-0 780
		3.05 uM 953.67 nM	10.280:
		298.02 nM	2.110 7.860
		93.13 nM	1,140
	1	29.10 nM	2.8201
		9.09 nM	4.150
		2.84 nM	5.590
59-0102	284.34		
59-0102		100.00 uM	-24.350
		31.25 uM	-11.140
		9.77 uM	63.540
		3.05luM	121.320
		953.87 nM	79.530
		298.02 nM	72.460
		93.13 InM 29.10 InM	66.290
		9.09 nM	45.690 27.260
		2.84 nM	42.330
		888.18 pM	33.430
59-0103	313.38		
	313.30	100.00 uM	-29.69
		31.25 uM	-29.53
		9.77 UM	-28.22
		3.05 luM	-27.72
		953.67 nM	-5.58
		298.02 nM	
		93.13 nM	54.15 170.95
		93.13 InM 29.10 InM	
		9.09 nM	
	-	2.84 nM	210.39
		2.84 InM 0.80 InM	200.41
		U.BUINM	114 55
59-0104	297.31		
		100.00 uM	-29.84
		31.25 uM	-26.72
		9 77 luM	-29.2
		3 05 luM	27 05
		953.67 InM	24.37
		298.02 nM	196 42
		93.13 nM	213.89

		29 10InM-	220.04
		9.09 nM	245.421
		2.84 nM	182.45:
		0 80lnM	119.55
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59-0105	267.29		
		100 00 uM	-25.72
		31.25 uM	-15.89
		9.77 uM	31.7
		3.05 uM	54.17
		953.67 nM	53.67
		298.02 nM	41.35
		93.13 nM	44.5
		29.10 nM	39.02
		9.09 nM	25.38
		2.84 nM	31.7
		0.80 nM	18.05
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59-0106	297.31		
	2.57.01	100.00 uM	-14.05
		31.25 uM	223.52
		9.77 uM	202.58
		3.05 uM	107.73
		953.67 InM	71,3
		298.02 InM	44.84
		93.13 InM	26.541
		29.10 nM	23 05
		9.09InM	27.87
	i	2.84 inM	12.23
		0.80 inM	11.41
		9.05.0m	11.4
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59-0107	332.38		i
	556.501	100.00 uM	48.55
		31.25 uM	22.87
	i-	9 77 uM	7.19
		3 05 luM	
		953 67 InM	0.65
	<del></del>	298.02 InM	
		93 13 nM	1.09
			1.091

		9.09(nMC) 8	11 11 32 11 11
			-2.62
		0.80 nM	-16.11
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59-0108	316.31		
		100.00 uM	227.73
		31.25 uM	96.02
		9.77 uM	58.57
		3.05 uM	37.23
		953.87 nM	18.94
		298.02 nM	25.68
		93.13 nM	4.8
		29.10 nM	2.62
		9.09 nM	-48
		2.84 nM	3.921
		0.80 nM	4.14
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59-0109	316.31		
		100.00 uM	43.12
<u>:</u>		31.25 uM	27.64
		9.77 uM	5.89
		3.05 UM	6.32
	1	953.67 nM	13.51
		298.02 nM	7.85
		93.13 nM	3.711
		29.10 nM	-3.27
		9.09 nM	5.01
		2.84 nM	-4.58
		0.80 nM	6.98
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59-0110	285.29		
		100.00 uM	65.11
		31 25 uM	
		9.77 uM	-35.27
		3.05 uM	25.26
		953.67 nM	27.01
		298.02 nM	15.24

		93 13 InMCh 2	F 140.68:2 / 4.4
		29.10InM	5.891
		9 09 nM	5 4 5
		2.84 nM	10.24
		0.80 nM	4 14
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59-0111	152.15		
		100.001uM	23.380
		31.25 uM	22.330
		9.77 uM	12.260
		3.05 uM	5.390
		953.67 nM 298.02 nM	2.190
		93.13 nM	1.230
		29.10 nM	6.350
		9.09 nM	4.350
		2.841nM	4.350
		D.BDinM	3.230)
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59-0112	149.19		
	146.161	100,00 uM	2.870
		31.25 uM	4.870
		9.77 uM	2.750
		3.05 uM	3.790
		953.87 nM	4.270
		298.02 nM	1.150
		93.13 nM	9.530
		29.10 nM 9.09 nM	0.920
		2.84 nM	12.900
		0.80 nM	2.990
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59-0113			
35-0113	274.37	400.00.44	
		100.001uM 31.251uM	22.010
		9.77 uM	7.500
		3.05 uM	3.070
		953.87 nM	-0.760
		298.02 nM	-4.690
		93.13 nM	-4.790
		29.10InM	5.090
		9.09 nM	0.150
		2.841nM 0.801nM	-0.250
		U-OUTHM	0 150

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59-0114	475.54			
		100.00 uM	52.030	_
		31.25 uM	36.120	_
		9.77 uM 3.05 uM	25.840	
		953.67 nM	16.670	
		298.02 nM	9.420	_
		93.13 nM	-1.080	
		29.10 nM	2.160	_
		9.09 mM 2.84 nM	-6.000	_
		0.80 nM	2.470	_
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59-0115	318.87	1.		
	318.07	100.00 uM	. 73.700	
		31.25 uM	2.770	
		9.77 uM	-10.430	_
		3.05 uM	-12.340	_
		953.67 InM	-13.750	_
		298.02 nM 93.13 nM	-13.960	_
		29.10 inM	-11.940 -9.830	_
		9.09 nM	-8.820	_
		2.84 nM	-0.950	_
		0.80 nM	-0.0501	_
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9-0116	269.30			
		100.00 uM	31 380	_
		31.25 uM 9 77 uM	109.060	
		3.05 uM	231.070	
		953 67 InM	132.020	_
		298.02 nM	75.820	-
		93.13 nM	53.250	_
		29.10 inM	47.500	
		9.09 nM 2.84 nM	39.440 42.170	_
		0.80 nM	31.180	_
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59-0117	268.38			
		100 00 uM	-68.520	_

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		31 25 tuM-	1 37 450pt /
		3.05 uM	111 630
		953.67 nM	64 340 4.740
		298.02 nM	-19.270
		93.13 nM	-26.660
		29 10 nM	-28.880
		9.09 nM	-22 880
		2.84 nM	41 300
		0.80 nM	-39.2201
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59-0118	313,38		
		100.00 uM	-67.170
		31.25 uM	-56.580
		9.77 uM	-58.060
		3.05 uM	-55.720
		953.87 nM	-48.200
		298.02 nM	-50.300
		93.13 nM	-33.310
		29.10 nM	-47.340
		9.09 nM	-49.310
		2.84 nM	-56.200
		0.80 nM	-57.310
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59-0119	314.34		1 1
	314,34	100.00 uM	167.500
		31.25 uM	-29.240
		9.77 uM	-29.240
		3.05 uM	-52.030
		953.67 inM	-52.030
		298.02 nM	-53.870
		93.13 nM	-38.110
		29.10 nM	-55.100
		9.09 nM	-52.270
		2.84 nM	-53.500
		0.80 nM	-43,650
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59-0120	504 49		
		100.00 uM	-82.790
		31.25 uM	-80.470
		9.77 uM	-66.800
		3.05 uM	-80.790
		953.67 InM	-54,240
		298.02 nM	-45.250
		93.13 nM	-50.680

		29.101nM	
	. ,	9 09 nM	50.300
		2.84 nM	-50.300
		0.80jnM	-90.300
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59-0121	245.29		
		100.00 uM	-79.6901
		31.25 uM	-75.5901
		9.77 uM	25.8501
		3.05 uM	94.850
		953.67 nM	43.910
		298.02 nM	-1.800
		93.13 nM	4.150
		29.10 nM	-22.050
		9.09 nM	-31.110
		2.84 nM	-26.760
		0.80 nM	-28.270
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59-0122	333,39		
		100.00 uM	-19.050
		31.25 UM	-12.080
		9.77 uM	-7.610
		3.05 luM	25.210
		953.67 nM	83.580
		298.02 nM	87.220
		93.13 nM	63.890
		29.10 nM	42.680
		9.09 nM	45.320
		2.84 nM	37.780
		0.80 nM	27.030
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59-0123	347.42		
		100.00 uM	34.430
		31.25 uM	34.710
		9.77 uM	38.620
		3.05 UM	55.100
		953.67 nM	51.900
		298.02 nM	41.410
		93.13 nM	29.970
		29.10 nM	73.760
		9.09 nM	17.120
		2.84 nM	13.480
		0.80 nM	1 190

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100,00 uM		דו /טר	U	
100.00 IAM				
31.25 IuM 51.500  9.77 IuM 14.5 800  3.05 IuM 135.530  9.93.87 IuM 224.590  9.93.87 IuM 224.590  244.02 IuM 134.650  9.3.13 IuM 134.650  9.3.13 IuM 91.800  2.2.10 IuM 51.500  0.8.00 IuM 53.000  0.8.00 IuM 53.000  0.8.00 IuM 51.400  9.77 IuM 54.120  1.3.125 IuM 67.500  9.77 IuM 54.120  1.3.125 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700  9.53.87 IuM 27.700	59-0124	350 44		
9.77 (uM 14.5 840) 3.05 (uM 13.5 830) 9.93.87 (nM 224.599) 9.93.13 (nM 13.4 840) 9.13.13 (nM 9.1 840) 9.13.13 (nM 9.3 80) 9.13.13 (nM 9.3 80) 9.13.13 (nM 9.3 80) 9.14 (nM 9.3 80) 9.15 (nM 9.3 80) 9.16 (nM 9.3 80) 9.17 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80) 9.18 (nM 9.3 80)	1			30.0401
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931.87 (nM 284 890)  2340.07 (nM 224 290)  9.31.31 (nM 134 850)  9.31.31 (nM 134 850)  9.31.31 (nM 9.3 80)  9.30 (nM 9.3 80)  1.30 (nM 9.3 80)  9.30 (nM 9.3 80)  9.31 (nM 9.3 80)  9.31 (nM 9.3 80)  9.31 (nM 9.3 80)  9.31 (nM 9.3 80)  9.31 (nM 9.3 80)  9.31 (nM 9.3 80)  9.31 (nM 9.3 80)  9.31 (nM 9.3 80)  9.30 (nM 15.70)  9.30 (nM 15.70)  9.30 (nM 15.70)				
99.01 M 224.300   99.10 M 19.840   99.10 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.840   90.01 M 19.00				
99.13   MM   134.840   29.10   MM   91.890   9.00   MM   60.390   2.84   MM   60.390   2.84   MM   60.390   3.08   MM   51.490   3.12   MM   51.490   3.12   MM   67.535   3.12   MM   67.535   3.12   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.13   MM   67.535   3.14   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15   MM   67.535   3.15				
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100.00 uM				
31.25 luM   67.530     9.77 luM   54.120     3.05 luM   28.700     3.05 luM   28.700     95.35 7 lnM   27.580     2280 2 lnM   22.280     93.13 lnM   22.700     23.10 lnM   18.500     9.09 lnM   18.700     2 84 lnM   9.840		372.45	100 00 144	4330
9.77/uM 54.120 1.030/uM 22.700 9.53.57/nM 21.590 29.02/nM 22.200 9.13/nM 22.200 9.13/nM 15.300 9.10/nM 15.300 9.09/nM 15.700 2.8/nM 9.8/nM				
3 .05 luM 22.700   95.55 FIAM 21.800   22.200   22.200   22.200   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.50   23.5				
953.57 ink   21.590    293.02 ink   22.220    93.13 ink   22.700    29.10 ink   15.30    5.09 ink   15.700    2 8 ink   9.500				
298.02 nM   22.280    93.13 nM   22.700    23.01nM   1.530    9.08 nM   15.700    2 84 nM   9.840				
93.13 inM 22,700   23.10 inM 1,530   9.09 inM 1,5700   2.84 inM 9,540   9.54 inM 9,540				
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		298.02		8.66	
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		<del>-</del>	3.051uM	-12.92	
			953.87 InM	-12.92	
		<del></del>	298.02 nM	-13.54	
			93.13 nM		
			29.10 nM	-3.65 -7.68	
			9.09 nM		
			2.84 nM	-6.18	
			0.80 nM	-9.97	
			U.80 mM	-2.81	
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59-0133		327.34			i
		1	100.00 uM	-16.04	
		1	31.25 UM	-16.91	
			9.77 uM	-17.31	7
			3.05 uM	-16.7	
		T	953.67 nM	-9.34	
			298.02 nM	-12.69	<del></del>
			93.13 nM	-11.23	<del></del>
		T	29.10 nM	-17.74	
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			2.84 inM	4 71	<del></del>
			0.80 nM	0.55	<del>- i</del>
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59-0135	356.39		1	1		1
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		31.25		-14.16		
		9.77		-1.98		
		3.05		0.97		
		953.67		11.68		-
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		100.001	uM	-6.91	
		31.25		-12.68	
		9.77		4.59	
		3.051		32.61	
		953.67		19.07	
		298.021		8.18	
		93.13		2.25	
		9.09		12.22	
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59-0139	340.43				1
		100.001	uM	45.53	
		31.251		44.59	
		9.77		53.62	
		3.051		30.42	
		953.67		26.25	
		298.021		20.311	
		93.131	nM	18.6	

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		9.09 nM	13.93	
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		31.25 µM 9.77 µM 3.05 µM 953.67 nM 298.02 nM 93.13 nM	5 69 19 85 43.96 44 73 37 12 24.36	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM	5 891 19 851 43 96 44 73 37 121 24 361 1 16.61	
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		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 uM 9.77 uM 3.05 uM 953.67 inM 298.02 inM 93.13 inM 29.10 inM 9.09 inM 2.84 inM	5 891 19 851 43.961 44.73 37 121 24.361 16.61 25.71	
		31.25 UM 3.05 UM 3.05 UM 3.05 UM 258.02 IM 258.02 IM 258.02 IM 258.02 IM 259.00 IM 29.00 IM 29.00 IM 20.00 IM 20.00 IM	5.69 19.85 4.3.96 4.4.73 37 12 2.4.39 19.5 19.5 7.87	
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	953.	67 nM	36.94	
		02InM	27.23	
		13InM	15.991	
		10 nM	19.27	
		09 nM	14.42	
		84 InM	11.33	
	0.	BO nM	23.72	
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59-0144	315.40	1		
	100.0	Mul00	-14.59	
		25 luM	441	
		77 uM	47.11	1
		25 uM	53.891	
		7 nM	43 11	-
		2 nM	29 21	
		3InM	18.51	
		0 nM	12.91	
		Mnie	5.54	
	, 2.8	MINM BOINM	3.71	
	0.8	MID	5.87	
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9-0145	350.27			i
	100.0	MulDi	435.91	
	31.2	5 uM	422.15	
		7 uM	446.93	1
	3.0	15 uM	434.17	-:
	953.6	7 nM	238.34-	
		2 nM	45.991	
	93.1	3 nM	9.22	
		0 nM	7.71	

	2.84 inM	6.271	
	2.04 INM	3.55	
	- Jackson	3.551	
59-0146	248.27		
	100.00 uM	-63.05	
	31.25 uM	4 42	
	9.77 uM	-13.73	
	3.05 uM	-16.45	
	953.67InM	-35.47	
	298.02 InM	-51.25	
	93.13 nM	-50.13	
	29.10 nM	-42.92	
	9.09 nM 2.84 nM	-45.64	
	2.84 inM 0.80 inM	-56.58 -39.68	
	U.SUINM	-39.68	
59-0147	314.36		
	100.00 luM	-85	
	31.25 uM	-85	
	9.77 uM	-80.29	
	3.05 iuM	-41.67	
	953.67 inM	78.691	
	298.02 nM	269.13	
	93.13 nM 29.10 nM	323.59	
	29.10 nM 9.09 nM	339.88	
	2.84 nM	270.46	
	0.80 InM	180.33	
	U.BU.IIM	180.33	
59-0148	291.35		
	100.001uM	-68.38	
	31.25 uM	-36.33	
	9.77 uM	-2.3	
	3.05 juM	12.12	
	953.67 InM	-2.42	
	298.02 nM	-16.21	
	93.13 nM 29.10 nM	-30.87	
	29.10 nM	-35.58	
	2.84inM	-39.07i	
	0.80 inM	45.53	
	J.301118		

59-0149	329.33		
	100.001uM		1
	31.25 uM	-16.9	
	9.77 uM	-0.53	
	3.05 uM	15.29	
	953.67 nM	78.78	
	298.02 nM	163.5	
	93.13 nM	223.57	
	29.10 nM	173.93	
	U.UU RM	122.3	
	2.84 INM	98.02	
	0.80InM	69.06	
59-0150	304.39		
	100.00 uM	63.32	
	31.25 UM	193.53	
	9.77 UM	419.26	
	3.05 uM 953.67 nM	497.21	
	298.02 InM	193.35	
	93.13 inM	99.46	
	29 10 inM	59.96	
	9.09 InM	59	
	2.84 nM	52.16	
	. 0 80 nM	48.751	
9-0151 9-0151	278.311		1
9-0101	100.00 uM	-6.660	
	31.25 uM	16.240	
	9.77 uM	18.300	
	3.05/uM	11.690	
	303.07 IIM	8.500	
	298.02 nM	9.070	
	298.02 nM 93.13 nM	9.070 6.110	
	298.02 nM 93.13 nM 29.10 nM	9.070 6.110 5.880	
	298.02 nM 93.13 nM 29.10 nM 9.09 nM	9.070 6.110	

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59-0152	266.275			
59-0152		100.00 uM	-6.8901	
		31.25 uM	12.490	
		9.77 uM	21.950	
		3.05 uM	12.820	
		953.67 nM	7.350	
		298.02 nM	4.290	
	+	93.13 nM	9.750	
		29.10 nM	4 860	
	+	9.09 nM	1.320	
	+	2.84 nM	4 280	
	<del>                                     </del>	0.80 nM	4.160	
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59-0153	282.73		1 1	-
59-0153		100.00 uM	-4.150)	
	1	31.25 JuM	-0.390	
		9.77 uM	11.120	
		3.05 uM	14.540	
		953.67 nM	9.520	
		298.02 nM	11.570	
		93.13 nM	-0.160	
		29.10 nM	1.550	
	+	9.09 nM	-0.960	
		2.84 nM	4.730	
	<del> </del>	0.80 nM	5.650	
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	262.312	100.00 luM	0.290	
	262.312	31 25 uM	24.670	
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	262.312	31 25 luM 9.77 luM 3.05 luM 953.67 lnM 298.02 lnM	24.670 15.680 14.540 13.170 5.540	
	262.312	31 25 JuM 9.77 JuM 3.05 JuM 953.67 JuM 298.02 JuM 93.13 JuM	24.670 15.680 14.540 13.170 5.540 2.690	
	262.312	31 25 iuM 9.77 iuM 3.05 iuM 953.67 inM 298.02 inM 93.13 inM 29.10 inM	24.670 15.880 14.540 13.170 5.540 2.690	
9-0154 9-0154		31 25 JuM 9.77 JuM 3.05 JuM 953.67 JuM 298.02 JuM 93.13 JuM	24.670 15.880 14.540 13.170 5.540 2.990	

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59-0155			1	1
59-0155	316.282	100.00 uM		
	1	31.251uM	1.900	
	1	9.77 uM	-9.450	
		3.05 uM	-0.220	
		953.87 nM	0.690	
		298.02 nM	5.090	
		93.13 nM	-3.250	1
	-	29.10 nM	0.530	
	<del>                                     </del>	9.09 nM 2.84 nM	-1.900	L. U.
	<del>                                     </del>	0.80 nM	9 480	
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59-0156 59-0156	333.391			
33-0136	-	100.001uM	5.840	
	-	31.25 luM	2.050	
	-	9.77 LuM 3.05 LuM	7.9601	1
		953.67 InM	-0.370	
	1	298.02 nM	-1.8801	
		93.13 nM	-3,550	
		29.10InM	-7.340	
		9.09 nM	-1 590	
		2.84 nM	2.850	
		0.80 nM	2.500	
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59-0157	290.366		i	1
59-0157	200.000	100.001uM	-6.440	
		31.25 uM	14 920	
		9.77 uM	19.930	
		3.051uM	11.440	0 -
		953.67 nM	6.570	
		298.02 nM	-7 190	
		93.13 nM	0.080	
		29.10 nM	-0.230	
		9 09 nM	-4.460	
		2.84 inM 0.60 inM	2.200	
		U BUINM	9.920	

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59-0158 59-0158	308.337				
39-0158		100.00		-5.980	
		31.25		3.720	
		9.77		16.140	
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		298.02		9.930	
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59-0159	308.337				
59-0159		100.00	uM .	2.790	
		31.25		13.530	
		9.77		4.700	
		3.05		10.910	
		953.67		2.800	
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9-0160		- 1		1 1	i
9-0160	319.408				
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		9.77		-3.390	
		3.05		5.300 15.910	
		953.67		6.610	
		298.02		11.380	
		93.13		4.460	
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		2.841		-0.650	
		0.80	1M	7.550	

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59-0196	323.201					İ
59-0196		100.00	uM	+	+	+
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9-0199	291.35				Ť	
9-0199	291.35	100.00	ıM			
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99-0204 344.389 190.00 luM 3 122 luM 3 20 luM 3 30 luM 953.87 luM 220.27 luM 230 luM 240 luM 250 luM			0.80	InM		7	+
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93-0215	182.957	
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33-0399	121 001	
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380.253	13.149 2.630	_
360.233	0.526	-
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3-0587	222.953 u	
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224.263	4.459	
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115,230 88,110 20,870 -28,680 5,250
128.130 38.560 41.240 -4.910 3.910
178.130 60.410 -0.180 -3.470 -8.490
-42.000 119.130 67.930 8.520

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93-1340	196 576	
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93-1474	145 940	
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-45.110 110.290 35.080 108.040 40.130	
75,940 173,150	

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850-7377	
350-7377	131 062 uM
	13.106
381,49	2 621
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50-7413	
50-7413	111.964 uM
	11.196
446.57	2.239
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50-7449	69.938 UM
	6.994
714.923	
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2.600 -7.350 -25.160
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-40.44 -2.55 157.01 78.73 23.91
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850-7485	143.096	Mu	-42.91
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850-7991	127.367		
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392.565	2.547	_	105.51
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850-8170	101.513	м	-33.79
	10.151	_	158.65
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850-8205		
850-8205	104 478 uM	-39 52
	10 448	51.18
478.57		163.82
	0.418	106.06
CHIRAL	0.084	73.58
500-8241		
850-8241	82.279 uM	-2.07
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607.685	1.646	118.23
	0.329	66.73
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850-8278		
850-8278	139.101 uM	40.09
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359.451	2.782	182.38
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850-8387	122.39	
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408.523		
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850-8469	87 921	u.
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568.692	1.758	
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-17 00 130.3 129.75 62.66 40.74	5
-21.13 11.30 131.92 71.13 58.56	
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850-8564	95.156	uM :		-30.92
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850-9071	109.998	M	[	-24.620
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850-9179		- 1		
850-9179	105.357	м	-2	4.6
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474.579	2.107		81	9.2
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GEN-9212	92.139 u	M	-26	
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542.657	1.843	_	111	
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850-9287	147 17	
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850-9356	99.506	uM
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896-0262	166.019 ul	4 -19.18
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895-0268	128.383 uA	
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389.458	12.638	40.25
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413.58	2.418	_
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95-0967	159.028	1184
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314.407	3.181	_
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895-1888	212.504	Mu
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895-2474	i	
895-2474	184.952	
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895-2475	162.159	M
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-33.66 29.75 148.84 73.77 28.14	
-20.74 128.69 56.37 43.27 19.44	
265.41 287.85 227.34 65.40 28.98	

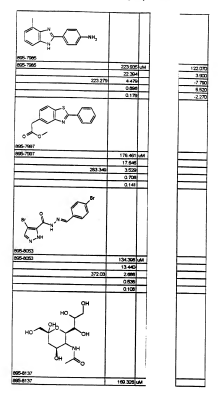
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283.331	3.529	283
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895-4843	159.581 uM	-17
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313.312	3.192	100
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896-5185	162.433 uM	-6
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895-7828	184.973	ωM
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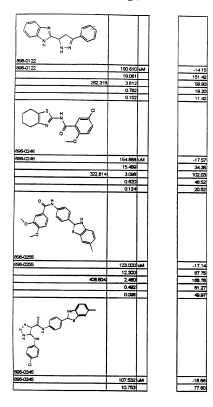


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895-9683	113.552	uM
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895-9898	178.349	uAI Na
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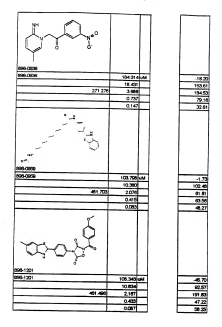


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464 979	2.151	188 94
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896-0390 896-0390	100 710 11	
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388.445	12.872	87 23
300.448	0.515	210.25
	0.103	73.36 28.25
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598-0506	ī	
896-0535	132.810 uM	-10.41
	13.281	73.84
376.478	2.656	199.80
	0.531	102.12
	0.106	35.72
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896-0554	121.499 uM	-16.32
	12.150	105.48
411.527	2.430	115.43
	0.466	53.86
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896-0692	424.000		
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380.897	2 625	-	149.23
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896-0719	91.950	м	-6.49
	9.195		187.43
543.774	1.839		127.43
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	0.074	_	36.16
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896-0773	147.228	M I	-13.94
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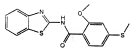
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896-0853	157 546	uM.
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317.367	3.151	
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896-0921	174.583	Mu
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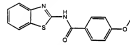


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350.266	0.571	

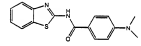
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-39.97 55.00 122.66	2
67.2s	
1,073.9 1,082.1 884.7	



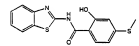
59-0072



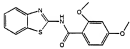
59-0102



59-0070



59-0144



59-0147

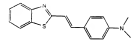
Max: 121 % EC50: 30 nM

Max: 214 % EC50: 200 nM

Max: 54 % EC50: 2 μM

Max: 340 % EC50: < 0.8 nM

EC50 : < 0.8 n

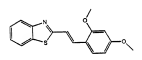


59-0099



Max: 269 % EC50: < 0.8 nM

Max: 200 % EC50: 30 nM



59-0210

FIG. 5B



59-0192 Max: 155 % EC50: 20 nM



59-0195 Max: 155 % EC50: 20 nM



59-0193 Max: 95 % EC50: 30 pM



59-0196 Inactive



59-0194 Inactive



59-0197 Max: 162 % EC50: 150 nM



59-0202 Max: 155 % EC50: 150 nM



59-0204 Max: 70 % EC50: 50 nM



59-0205 Max: 250 % EC50: < 0.8 nM

59-0206 Max: 150 % EC50: 20 nM

59-0207 Max: 50 % EC50: 100 nM

59-0208 Max: 85 % EC50: 1 uM

50-0197 Max: 245 % EC50: 3 nM

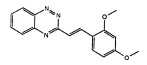
59-0078 Max: 380 % EC50: 1 nM

FIG. 6A

59-0199 Max: 170 % EC50: 100 nM

59-0203 Max: 275 % EC50: < 1 nM

59-0286 Max: 160 % EC50: 300 nM



59-0285 Max: 200 % EC50: 30 nM

FIG. 6B

R =



59-0030 Max: 90 % EC50: 1 uM

59-0089 Max: 120 % EC50: 5 uM

59-0093 Max: 35 %

59-0094 Max: 45 %

59-0091 Max: 96 % EC50: 1 uM

59-0090

59-0150 Max: 500 % EC50: 1 nM

Max: 41 %

59-0199 Max: 170 % EC50: 100 nM

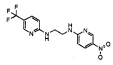
59-0092 Max: 50 % EC50: 10 uM

59-0198 Max: 135 % EC50: 100 nM

FIG.

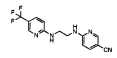
59-0145

Max: 300 % EC50: 0.5 uM



59-0450

Max: 270 % EC50: 5 uM



59-0483

Max: 260 % EC50: 3 uM

59-0459

Max: 180 % EC50: 5 uM

59-0480

Max: 180 % EC50: 5 uM

FIG.

FIG. 8 #

59-0098 FIG.

$$\begin{array}{ll} \text{Max}: & 48~\% \\ \text{EC50}: & 30~\mu\text{M} \end{array}$$

Max: 222 % EC50: 20 nM

83

X, Y = F, Cl, OMe < 50 % max @ 100 uM

### 59-0098 Analogs

X, Y = F, Cl, OMe < 50 % max @ 100 uM

### 59-0096 Analogs

X, Y = F, Cl, OMe < 50 % max @ 100 uM

#### 59-0097 Analogs

8C

FIG.

_		

	Compound		Max Response of	ZGI Score in Ex Vivo	OS Sereen in Ex Vivo
Compoun		EC50	59-0008	Assay	Assay
				7.0001	Hoody
59-0364	Р	0	0	1	
59-0076	Р	0	م ا	1	
59-0451	P	0	ō	i	
59-0472	P	Ó	Ö	i i	
59-0073	Р	o	o i	'	1+
59-0095	н	??	0.5x (30 uM)		1
59-0471	Р	??	0.5x (100 uM)	1	'
59-0030	a l	??	.7x ( 1uM)	i i	1,1+
59-0470	P	50 uM	1.2x (100 uM)		1,17
59-0450	Р	5 uM	2.7x (30 uM)	' '	
59-0459	P	5 uM	2x (10 uM)	1	
59-0064	a	3 uM	1.5x (? uM)	<u>i</u>	

59-0008	Q	1 uM			1
59E0115	Cost This is	2(6(6)) (1)	4x (SalMi)	STATE OF THE PARTY	100 (100) Table
59-0106	T	300 nM	2x (9 uM)	NAME OF TAXABLE PARTY.	1
59-0070	T	200nM	2x (3 uM)		1,1+
59-0097	н	100 nM?	2x (30 uM)		1+
59-0096	Н	100 nM?	4x (100 uM)		1 1
59-0116	н	30 nM	2.5x (3 uM)		1+,2-
59-0210	T	30 nM	2x (3 uM)		1
59:0098	10 - 12 ( - 1)	Per chiye	28 9 UM	Page.	Company Company
59-0019	Q	10 nM	2.5x (300 nM)	1+.2-	1.1+
59-0078	Q	9 nM	4x (1 uM)	•	1
59-0045	H	5 nM	4x (1uM)	1	1
50-0197	Q	3 nM	2.5x (300 nM)	1	1+,2-
59-0099	T	2 nM?	3x (1 uM)		1,1+
59-0282	Q	1 nM	2x (3 uM)	2	1+,2-
5880208	Section .	1 11/4	Service IMI	A STATE OF THE STA	4.3
59-0072	T	300 pM	2x (uM)	1-1+	1.1+
59-0150	Q	<1 nM	5x (3 uM)	1-2?	1
59-0104	Т	<1 nM	2x (uM)	1+.2-	1
59-0103	Т	<1 nM	2x (30 nM)		1,1+
59-0124	Т	<1 nM	2.5x (1 uM)		1+,2-
59-0205	T	<1 nM	2x (2 nM)		i

H = Hydrazone/Hydrazide (45) Q = Quinoline/Quinoxaline (197) P = Bis-pyridines (145)

T = Benzothiazole (104)

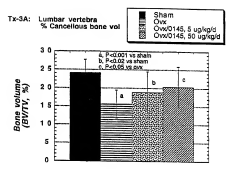
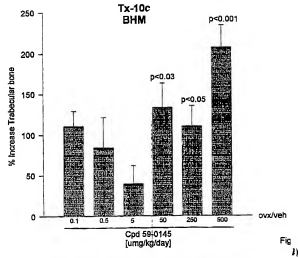


Fig 10



% Increase of trabecular bone over the ovx/vehicle group

Tx-10c

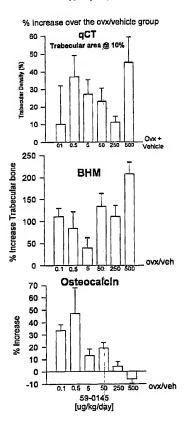


Fig 12

MOLSTRUCTURE	MOL>NNCIMOL WEIGHT NUM1			
٥١٠٥.	59-0020	268.732		
CO-S	59-0021	284.723		
a a	59-0022	266.367		
8	59-0023	239.276		
	59-0008	254.315		
	59-0024	220.276		
arar.	59-0025	224.308		
	59-0026	248.29		
ಯ್ಯಂ	59-0027	250.303		
مرم	59-0028	226.283		
مئنی	59-0029	249.272		

Figure 13

59-0031	231.3	
	1	
1 ,		
59-0030	233,275	
59-0032	248.287	
59-0033	248.287	
59-0034	268.343	
59-0035	291.356	
59-0036	262.314	
59-0037	30В	
59-0038	241.295	
59-0039	312.352	
59-0040	290.368	
59-0041	501.902	
	59-0032 59-0033 59-0034 59-0035 59-0037 59-0039 59-0040	59-0032 248.287  59-0033 248.287  59-0034 268.343  59-0035 291.356  59-0036 262.314  59-0037 308  59-0038 241.295  59-0039 312.352

Page 2

59-0042	281.36	
59-0043	260.288	
59-0044	341.21	
59-0045	283.333	
59-0046	389.372	· · · · · · ·
59-0047	303.367	
59-0048	384.501	
59-0049	251.29	
59-0050	303.364	
59-0051	251.353	
59-0052	393.276	
59-0053	354.412	
	59-0043 59-0044 59-0045 59-0046 59-0048 59-0050 59-0050	59-0043 280.288 59-0044 341.21 59-0045 283.333 59-0046 389.372 39-0047 303.367 59-0049 251.29 59-0050 303.364 59-0051 251.353 59-0052 393.276

مثن	59-0054	236.276	
36 7	59-0055	425.508	
ith.	59-0056	512.341	
CT's->-Con,	59-0102	284.339	
CT, C.	59-0057	329,448	
mas Color	59-0058	268.34	
	59-0059	375.923	 
8-4-	59-0060	301.391	
	59-0061	255.3	
g-100	59-0062	357.44	
	59-0063	255.344	
000	59-0064	276.385	

OH S	59-0065	254.313	
	59-0066	248.33	
Cin, in	59-0067	254.315	
	59-0068	259.354	
HO CONTRACTOR	59-0069	268.223	
0,0	59-0019	275.353	
CH, CH,	59-0070	297.38	
من م	59-0071	291,352	
	59-0072	330.431	
⁶⁰⁰⁰ k	59-0073	376,303	
-35 +0,-55+	59-0074	642.735	
<del></del>	59-0075	618.775	

, _a a .	59-0076	463.208	
+			
XXCODY	59-0077	445.193	
CO-Oz-	59-0078	276.341	
009	59-0079	231.297	
	59-0080	284.338	
	59-0081	377.466	
CY a cris	59-0082	222.267	
ಹ್ಹೆ	59-0083	330.414	
Cho.	59-0084	264.283	
	59-0085	278.31	
and.	59-0086	292,293	
coira-	59-0087	291.309	
· · · · · · · · · · · · · · · · · · ·			

^^	59-0088	263.299	
W.J. O.		200.255	
	59-0089	281.357	
and.	59-0090	324.425	
0000	59-0091	307.394	
₩ ₩	59-0092	281.357	
00-0	59-0093	232,285	
	59-0094	282.345	
2,02	59-0095	299.328	
graf	59-0096	313.355	
grat	59-0097	330.41	
المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات المرات ال	59-0098	325,368	
C:\\Q_\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	59-0099	280.393	

	59-0100	254.719	
	59-0101	230.232	
	59-0103	313,379	
CT has prom	59-0104	297.312	
CT Co.ch.	59-0105	267.287	
	59-0106	297.312	
ي الم	59-0107	332,378	
2000	59-0108	316,311	
	59-0109	316.311	
5,000	59-0110	286.286	
H _M N J ¹ OH	59-0111	152.152	
O CH ₃	59-0112	149.192	

~~	59-0113	274.365	
		274.365	
"000 B"	59-0114	475.548	
~~~~	59-0115	318.87	
Dyr Oyon	59-0116	269.302	
H ₂ C CH ₃	59-0117	268.352	
وثث	59-0118	313.354	
N,C O CH	59-0119	314.335	
1985-tsk <u>i</u>	59-0120	504.485	
CITO CITO	59-0121	245.284	
-During	59-0122	333.389	
مثيه	59-0123	347.416	
Jamo	59-0124	350,44	

60°	59-0125	372.447	
٥٥٥	59-0126	260.295	
To the second	59-0127	329,405	
mig-	59-0128	436.34	
9	59-0129	277.713	
	59-0130	287.345	
CHA.	59-0131	331.225	
8 /	59-0132	313.315	
0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	59-0133	327.342	
) 3 km 3 km	59-0134	357.367	
88	59-0135	356.383	
	59-0136	411.868	

0.	59-0137	296.712	
		298.712	
2,00	59-0138	340.808	
ಭಿಂ	59-0139	340.424	
000	59-0140	289.164	
16.89 \$-	59-0141	437.324	
24	59-0142	379.288	
S	59-0148	447.285	
CTS HO SCH,	59-0144	316.404	
400-04	59-0145	350.265	
	59-0146	246.268	
as to	59-0147	314.364	
and	59-0148	291.352	

59-0150 59-0151	304.391	
59-0151		
	278.31	
59-0152	266,274	
59-0153	282.729	
59-0154	262.311	
59-0155	316,281	
59-0156	333.389	
59-0157	290.364	
59-0158	308.335	
59-0159	308.335	
59-0160	319,406	
	59-0153 59-0154 59-0155 59-0156 59-0157 59-0158	59-0154 282,729 59-0155 316,281 59-0156 333,389 59-0157 290,384 59-0158 308,335 59-0159 308,335

CO. C.	5 9-0 161	291.352	
	59-0162	287.321	
W.C	59-0163	249.272	
aj.a	59-0164	299.332	
CO TO	59-0165	250.26	
COTO	59-0166	270.334	
	59-0167	263,299	
	59-0168	269.346	
(C)-(C)	59-0169	288.309	
CO. TO	59-0170	250.26	
CC), ID	59-0171	238.249	
arian	59-0172	306.32	

	59-0173	299.3321	
		299.332	
The Con	59-0174	279.298	
W. C.	59-0175	306.348	
	59-0176	256.288	
CL, TO	59-0177	251.248	
	59-0178	239.237	
arto.	59-0179	257.292	
arara.	59-0180	417.487	
	59-0181	313,358	
CT TO	59-0182	288,309	
مانی	59-0183	305.36	
07,0	59-0184	252.272	

	59-0185		
	59-0185	345.444	
CIS NO SE	59-0186	374.382	
0,38%	59-0187	389,494	
ze exp.	59-0188	616.784	
\$20~	59-0189	490.579	
9, 10, 10, 10, 10, 10, 10, 10, 10, 10, 10	59-0190	550.631	
asto.	59-0191	584,605	
	59-0192	344.389	
	59-0193	344.389	
	59-0194	344.389	
	59-0195	318,783	
المراجعة الم	59-0196	323,202	

CT S N CT CI	59-0197	323,202	
Chara Con	59-0198	261.323	
	59-0199	291.348	
guá	59-0200	342.349	
£ 145.	59-0201	331.326	
	59-0202	300,337	
	59-0203	292.336	
\$\\phi_{\phi}	59-0204	344.389	
CT STO CH,	59-0205	318.783	
The same	59-0206	348.809	
المرابع المراب	59-0207	348.809	
a last	59-0208	338.308	

Page 16

	155 555		
a la la la la la la la la la la la la la	59-0209	247.296	
\$\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	59-0210	297.376	
CH _s	59-0211	264.926	
CH ₈	59-0212	314.364	
	59-0213	294.333	
	59-0214	348.809	
مهج	59-0215	340.401	
Character of the contract of t	59-0216	264.304	
**************************************	59-0217	278.331	
275.	59-0218	292.357	
er in	59-0219	329.379	
	59-0220	300.312	

-o CM	59-0221	283,329	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	59-0222	309,367	
**************************************	59-0223	284.27	
d.gr	59-0224	330,338	
MO CO OH	59-0225	256.26	
" J. " J'o	59-0226	285,258	
\$ 3	59-0227	296,396	
CH ₉ CH ₉	59-0228	269.846	
CH ₉	59-0229	239.32	
ON-O.	59-0230	284.317	
Have I Cha	59-0231	318,399	
H ₁ C CH ₀	59-0232	269.35	

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A .N	59-0233		
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Ch-75-"	59-0236	280.325	
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	59-0238	340.401	
ans.	59-0239	330.338	
Erg.	59-0240	347.393	
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3400	59-0261	364.423	
	59-0262	398.36	
	59-0263	368,455	
	59-0264	383.254	
	59-0265	393.26	
	59-0266	328,39	
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Typia	59-0271	360.364	
Eng.	59-0272	381,838	
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	59-0276	358.273	
	59-0279	327.406	
Z, SS-	59-0277	372.375	
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	59-0310	368,378	
	59-0311	287.705	
	59-0313	293.127	
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	59-0315	275.137	
M.C. T. C.	59-0316	303.191	
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7770	59-0318	326.679	
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000	59-0320	206.247	
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H.C. C.H.	59-0322	284.745	
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	59-0312	309.582	
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	59-0368	311.339	
96	59-0369	387.437	
	59-0370	328.39	
	59-0371	372.399	
۵۶-۶۶۰۶۳ (59-0372	399.469	
	59-0373	299.353	
	59-0374	255.363	
م م	59-0375	261.391	
	59-0376	331.351	
,	59-0377	351.408	
	59-0378	285.389	
Cit.	59-0379	337.379	

Page 30

*contr	59-0380	408.813	
*anda	59-0381	408.813	
*ania	59-0382	408.813	
Thing	59-0383	468.699	
£	59-0384	340.405	
3.33	59-0385	334.377	
yar-oo	59-0386	367.761	
XXXX	59-0387	323.729	
ATT.	59-0388	451.23	
xanja.	59-0389	474.268	
* The state of the	59-0390	487.284	
****	59-0391	466.245	

the state of the s	59-0392	442.78	
*ar-ja	59-0393	395.767	
Harry C.	59-0394	393.195	
PACAL CAR	59-0395	370.804	
	59-0396	378.18	
Struck,	59-0297	424.808	
× ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	59-0398	414.234	
y though	59-0399	502.245	
*arfa	59-0400	526.388	
	59-0401	364.197	
	59-0402	362,181	
HING.	59-0403	538.803	

TOTO.	59-0404	549.378	
	59-0405	437.315	
	59-0406	406.233	
×4.	59-0407	349.699	
de de la contraction de la con	59-0408	561.868	
in in the second	59-0409	535.821	
S. Co	59-0410	340.428	
Harton Confirm	59-0411	464.294	
	59-0412	429.849	
Har J. Co.	59-0413	459.874	
in ha	59-0414	497.846	
Stark	59-0415	518.905	

	59-0416	454.834	
State for an	59-0417	484.86	
940	59-0418	333,268	
O)~;;;.	59-0419	367.761	
Y TOTAL	59-0420	352.767	
*axig	59-0421	539.339	
مثم	59-0422	351.253	
å m	59-0423	385.698	
*,C.**	59-0424	494.186	
*X-2-	59-0425	400.186	
* The state of the	59-0426	380.756	
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59-0428	380.756	
59-0429	409.793	
59-0430	313.669	
59-0431	454.859	
59-0432	395.767	
59-0433	407.821	
59-0435	433.738	
59-0436	444.637	
r		
59-0439	525.826	
	59-0430 59-0431 59-0432 59-0433 59-0435	59-0429 409.793 59-0430 313.669 59-0431 454.859 59-0432 395.767 59-0433 407.821 59-0435 433.736 59-0436 444.637

59-0440	525.8261	
59-0441	311,339	
59-0442	303.704	
59-0443	387,256	
59-0444	269,259	
	404.356	
59-0446	404.356	
59-0447	352.241	
59-0448	314.39	
	394.274	
59-0450	329,281	
59-0451	384.71	
	59-0442 59-0443 59-0444 59-0445 59-0446 59-0447 59-0449 59-0449	59-0442 303.704 59-0443 337.256 59-0444 269.259 59-0445 404.356 59-0446 404.356 59-0447 352.241 59-0448 314.39 59-0449 394.274 59-0450 329.281

Page 36

	59-0452	242,324	
0.00	59-0453	214.271	
Janes .	59-0454	264.291	
<u>z</u> 5	59-0455	300.32	
-12-02-	59-0456	308.296	
"-ioo	59-0457	330.342	
C*C**	59-0458	300.408	
*aca;	59-0459	364.292	
×a~o	59-0460	252.238	
	59-0461	266.265	
*Q~~Q	59-0462	280.292	
×a~o	59-0463	253.226	

. f	59-0464	267.253	
yana,	59-0465	363,26	
	59-0466	315.352	
	59-0467	212.294	<u>-</u>
<u></u>	59-0468	213,283	
	59-0469	378,318	
OO	39-0409	376,316	
	59-0470	325.293	
40~0×	59-0471	350.261	
40-0g	59-0472	351.249	

or po	59-0476	350.265	
4000	59-0477	283,256	
*ang	59-0478	351.253	
مسمر	59-0479	283.258	
yours	59-0480	332.328	
yarak	59-0481	363.26	
x0~~0+	59-0462	349.277	
B-DK	59-0483	307.278	
25	59-0484	315.246	
Q~~5	59-0485	250,3	
*ariax	59-0486	364,292	
arak	59-0487	302.298	

59-0488	486.259	
59-0489	255.3	
59-0490	322.909	
59-0491	317.269	-
59-0492	289.161	
59-0493	364.248	
59-0494	232.285	
59-0495	299.294	
59-0496	354.33	
59-0497	340,303	
59-0498	282.268	
59-0499	296.294	
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International application No. PCT/US97/18864

 CLASSIFICATION 	OF SUBJECT	MATTER
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IPC(6) :Please See Extra Sheet. US CL :Please See Extra Sheet.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : Please See Extra Sheet.

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

CAS--structure APS--diaryl, bone, osteo?, BMP

DIALOG-diaryl, bone, osteo?, BMP

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	US 5,441,964 A (BRYANT et al.) 15 August 1995, see entire document.	1-2, 5-28, 55-56
Y	US 5,523,309 A (BRYANT et al.) 04 June 1996, see entire document, especially claim 8.	1-2, 5-28, 55-56
Y,P	US 5,622,974 A (MUEHL) 22 April 1997, see entire document, especially claim 5.	1-2, 5-28, 55-56
Y	WO 93/10113 A1 (TEIKOKU HORMONE MFG. CO., LTD.) 27 May 1993, see entire document.	1-2, 5-28, 55-56
Y	WO 95/10513 A1 (PFIZER INC.) 20 April 1995, see entire document, especially claim 20.	1-2, 5-30, 55-56
Y	US 5,280,040 $$ A (LABROO et al.) 18 January 1994, see entire document.	1-4, 31-43, 55-56

х	Further documents are listed in the continuation of Box (с. 🔲	See patent family annex.	
	Special categories of cited documents:	*T*	later document published after the international filing date or priority	
۰۸*	document defining the general state of the art which is not considered to be of particular relevance		date and not in conflict with the application but cited to understand the principle or theory underlying the invention	
ъ.	earlier document published on or after the internstional filing date	•x•	document of particular relevance; the claimed invention cannot be	
1,*	document which may throw doubts on priority claim(s) or which is		when the document is taken alone	
	cited to establish the publication date of another citation or other special reason (as specified)	•Y•	document of particular relevance; the claimed invention cannot be considered to involve an inventive stap when the document is	
•0•	document referring to an oral disclosure, use, exhibition or other means		combined with one or more other such documents, such combination being obvious to a person skilled in the art	
P.	document published prior to the international filing date but later than the priority date claimed	.w.	document member of the same patent family	
Date o	of the actual completion of the international search	Date of	mailing of the international search report	
28 J	ANUARY 1998	26	FEB 1998	
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International application No. PCT/US97/18864

C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	Chem. abstr. Vol. 127, abstract No. 127:17703, PETRIE et al. Preparation of (hetero) aromatic compounds for treating bone deficit conditions', WO-97/15308 (Eng.).	1-4, 31-43, 55-56
Y	Chem. abstr. Vol. 107, abst. No. 107:109578, WATTS et al. 'Studies on the ligand specificity and potential identity of microsomal antiestrogen-binding sites', Mol. Pharmocol. 1987, 31(5), 541-51.	1-2, 50-56
Y	Chem. abstr. Vol. 108, abstract No. 108:69162, JORDAN et al. Effects of antiestrogens on bone in castrated and intact female rats', Breast Cancer Res. Treat. 1987, 10(1), 31-5.	1-2, 50-56
Y	Chem. abstr. Vol. 115, abstract No. 115:8533, SCHWARZ et al. 1,2-diphenyl-1-pyridybut-1-enes - potential antiestrogens. part 1. synthesis' Arch. Pharm. 1991, 324(4), 223-9.	1-2, 44-49, 55-56
Y	NEELAM et al. Structure-activity relationship of antiestrogens: A study using triarylbutenone, benzofuran and triayrlfuran analogues as models for triarylethylenes and triarylpropenones. J. Med. chem. 1989, Vol. 32, pages 1700-1707, see entire article.	1-2, 50-56
Y	VON ANGERER et al. Studies on heterocycle-based pure estrogen antagonists. Ann. N. Y. Academy Sciences. 1995, Vol. 761, pages 176-191, see especially pages 178-180.	1-2, 5-28, 55-56

International a	pplication	No
PCT/US97/18864		

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)		
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:		
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:		
Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:		
Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).		
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)		
This International Searching Authority found multiple inventions in this international application, as follows:		
Please See Extra Sheet.		
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.		
As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.		
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:		
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims, it is covered by claims Nos.:		
Remark on Protest		
No protest accompanied the payment of additional search fees.		

International application No.

PCT/US97/18864

A. CLASSIFICATION OF SUBJECT MATTER:

IPC (6): A61K 31/165, 31/215, 31/33, 31/405, 31/415, 31/42, 31/425, 31/44, 31/47, 31/505, 31/53, 31/535, 31/54

A. CLASSIFICATION OF SUBJECT MATTER:

US CL: 514/222.5, 223.2, 223.8, 224.2, 226.5, 229.2, 230.5, 255, 258, 259, 296, 307, 311, 336, 345, 352, 354, 457, 365, 367, 374, 375, 385, 394, 396, 397, 415, 443, 535, 646

B. FIELDS SEARCHED

Minimum documentation searched

Classification System; U.S.

514/222.5, 223.2, 223.8, 224.2, 226.5, 229.2, 230.5, 255, 258, 259, 296, 307, 311, 336, 345, 352, 354, 457, 365, 367, 374, 375, 385, 394, 396, 397, 415, 443, 535, 646

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING

This ISA found multiple inventions as follows:

This application contains claims directed to more than one species of the generic invention. These species are deemed to lack Unity of invention because they are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for more than one species to be searched, the appropriate additional search fees must be paid. The claims are deemed to correspond to the species as listed in the following manner.

Group 1, claims 3-4 and 31-43 compounds corresponding to Ar1 is condensed six membered heterocyclic ring, Ar2 is various aromatic rines:

Group II, claims 5-28, compounds corresponding to Ar1 is condensed five membered heterocyclic ring, Ar2 is various aromatic rings;

Group III, claims 29-30, compounds corresponding to Ar1 is isolated five membered heterocyclic ring, Ar2 is various aromatic rings;

Group IV, claims 44-49, compounds corresponding to Ar1 is isolated six membered heterocyclic ring, Ar2 is various aromatic rings;

Group V, claims 50-54, compounds corresponding to Ar1 is phenyl ring, Ar2 is various aromatic rings;

Group IV, claims 1-2, 55-56 in part (remaining compounds)

The following claims are generic: 1-2, 55-56

The species listed above do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2 and ANNEX B section (f), the species lack the same or corresponding special technical features for the following reasons:

The six groups of compounds corresponding to method of treating conditions of deficiency in bone growth, recoption or replacement using streaturally distinctive compounds. Each group of compounds as delineated above does not share significant structural element (see Ar1, Ar2 and L are all variables, thus, not common element). In addition, at least one Markush alternative is found in CA 127:17703.